Exhibit C

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE (Case No. 98,385-A)

Start PATENT

In application of: Hauptmann et al.) Before the Examiner: G. Draper
Serial No.: 08484,312) Group Art Unit: 1646
Filed: June 7, 1995)	
For: TNF RECEPTORS, TNF BINDING PROTEINS AND DNAs CODING)))
FOR THEM)

DECLARATION PURSUANT TO 37 C.F.R SECTION 1.132

- I, John D. Mountz, M.D. Ph.D., residing at 2800 Vestavia Forest Place, Birmingham, Alabama, United States of America, hereby declare:
- 1. I am Professor of Medicine in the Department of Medicine, Division of Clinical Immunology and Rheumatology, The University of Alabama at Birmingham. My Curriculum Vitae is attached hereto as Appendix A.
- 2. I have read U.S. Patent No. 5,695,953 (hereinafter, "the '953 patent"; attached hereto as **Appendix B**) and have considered the written description of the material disclosed in the patent and the extent to which this disclosure would enable one of ordinary skill to make and use recombinant DNA encoding a soluble tumor necrosis factor (TNF) inhibitor protein (hereinafter "TNF-IP").

Extent of Actual Experimental Disclosure in '953 Patent

- 3. The '953 patent discloses the existence of a TNF "inhibiting" activity in human urine (col. 4, ln. 9-10). This activity was associated with a protein having a molecular weight of 40-80 kilodaltons (kD) as determined by gel filtration chromatography (col. 4, ln. 10-13), termed TNF-IP.
- 4. The '953 patent describes generally a biochemical separation protocol for obtaining TNF-IP from urine of healthy individuals (col. 5, ln. 14-15). The biochemical separation protocol included membrane filtration to concentrate the urinary fluid (col. 8, ln. 30-37); two different cation-exchange chromatography steps (col. 8, ln. 40-59 and col. 8, ln. 61

- through col. 9, ln. 15), anion-exchange chromatography (col. 9, ln. 16-35), and reversed-phase high pressure liquid chromatography (col. 9, ln. 36-64).
- 5. The protein product of this biochemical separation protocol was analyzed by SDS-polyacrylamide gel electrophoresis (col. 9, ln. 65 through col. 10, ln. 19) and yielded a band migrating at a molecular weight of about 26-28 kD (col. 10, ln. 17-19), and further described as being 27 kD in size (col. 10, ln. 37).
- 6. This putative inhibitor protein was able to inhibit the biological activity of TNFα as demonstrated by treating TNF-sensitive cells with TNFα in the presence of the inhibitor protein (col. 6, ln. 24-40 (Table I); col. 6, ln. 52 through col. 6, ln. 57).
- 7. This protein product was reported to be over 40% "pure" and was also described as being the "major" protein in the preparation (col. 10, ln. 35-37).
- 8. From this partially-purified protein preparation, an amino acid sequence was obtained that was represented as being fourteen of the first sixteen N-terminal amino acids of TNF-IP, with one residue being identified as theoretical and a second residue being unidentified (col. 10, ln. 21-56).
- 9. This is the extent of the actual disclosure of the '953 patent, *i.e.*, what was actually done by the inventors.

Prophetic Disclosure in '953 Patent

- 10. In addition to the actual disclosure, the '953 patent contained the prophetic description of additional experiments directed toward isolation of a recombinant DNA molecule encoding the TNF-IP product of the biochemical separation protocol (col. 10, ln. 57 through col. 16, ln. 10).
- 11. This disclosure concerned experiments that were proposed but never performed by the named inventors of the '953 patent.
- 12. The '953 patent speculates that cloning can be achieved by conventional techniques, including describing in the most general terms three proposed methods for isolating TNF-IP encoding cDNA: using antibodies to screen λ gt11 cDNA libraries, using oligonucleotide probes to screen cDNA libraries or using oligonucleotide probes to screen genomic libraries.
- 13. The first two of these methods were absolutely dependent on identifying a cultured cell line expressing TNF-IP and preparing a cDNA library from such a cultured cell line

- (col. 11, ln. 40 through col. 12, ln. 5). The '953 patent predicted that a cell line expressing TNF-IP could be found by either immunofluorescence detection or "Western" blotting of candidate cell lines using an antibody prepared from TNF-IP (col. 11, ln. 29).
- 14. The '953 patent recommended that such an antibody be prepared by injecting rabbits or mice with either the protein product of the biochemical separation protocol described in the patent, or by using a synthetic peptide corresponding to the N-terminal amino acid sequence (as reported at col. 10, ln. 43) (col. 11, ln. 17-24). In a suggested alternative, the '953 patent teaches that the putative TNF-IP N-terminal peptide fragment could be used to produce a fusion protein in *E. coli* that could be purified and injected into mice (col. 11, ln. 19-22).
- 15. Once a cell line expressing TNF-IP had been identified using this scheme, the '953 patent taught that a cDNA library would be prepared from cellular mRNA using conventional techniques (col. 11, ln. 39).
- 16. In the first method suggested by the '953 patent, TNF-IP cDNA would be cloned into a λgt11 cloning vector, which enables expression selection of desired clones (col. 11, ln. 42). The supposed advantage of using this vector was that an antibody, such as antibodies produced according to the teachings of the '953 specification, could be used to screen the cDNA library.
- 17. In the second method, the '953 patent taught that a cDNA library could be screened with probes produced from the predicted nucleotide sequence of the putative N-terminal peptide of the putative disclosed TNF-IP protein (col. 12, ln. 5-15).
- 18. The '953 patent also taught that additional peptide sequences could be obtained from the putative TNF-IP by proteolytic fragmentation using known proteases (col. 12, ln. 19).
- 19. The '953 patent taught that it may be possible to employ an unfractionated set of oligonucleotides in the same manner in which one would employ a single oligonucleotide to clone the gene that encodes the TNF-IP. The use of such oligonucleotide or set of oligonucleotides containing the theoretical "most probable" sequence capable of encoding the TNF Inhibitory Protein gene fragments (following the "codon usage rules" disclosed by Lathe et al., 1985, J. Molec. Biol. 183: 1) permits one to identify the sequence of a complementary oligonucleotide or set of oligonucleotides which may be used as a probe for the gene of TNF-IP (col. 12, ln. 47 to col. 13, ln. 2).

- 20. The last of the three suggested methods disclosed in the '953 patent for isolating a DNA molecule encoding TNF-IP involved screening a library prepared from human genomic DNA with the same oligonucleotide probes as taught for screening cDNA libraries (col. 11, ln. 63 et seq.).
- 21. There was no disclosure of the results of any of the cloning experiments described in the '953 patent, and no evidence that any of these experiments had actually been performed at the time the application that matured into the '953 patent was filed.
- 22. Neither the nucleotide sequence nor production of any oligonucleotide, degenerate oligonucleotide or mixture of degenerate oligonucleotides was disclosed in the '953 patent.
 - 23. The identity of no cell line expressing TNF-IP was disclosed in the '953 patent.
- 24. The screening of no cDNA or genomic DNA library was disclosed in the '953 patent, and no DNA molecule encoding TNF-IP was disclosed in the '953 patent.

Failure of Inventors to Successfully Clone DNA by Prophetic Teachings

- 25. In a series of patent publications and other publicly-available references, some or all of the inventors named in the '953 patent reported that several of their attempts to isolate a DNA molecule encoding TNF-IP according to the methods described in the '953 patent were unsuccessful.
- 26. For example, Israeli Patent Application No. IL 92697 (Appendix C) discloses the applicant's failure to obtain a cDNA encoding TNF-IP by screening oligo dT-primed cDNA libraries prepared from human liver and placental tissue, or the human cell lines HeLa and U937 using a mixture of degenerate oligonucleotides prepared as described in the '953 patent and derived from the putative N-terminal amino acid sequence of the putative TNF-IP protein described in the '953 patent.
- 27. In this application, presumably at least 150 positively-hybridizing clones were obtained in an initial screening. Of these clones, three survived additional rounds of screening by positively-hybridizing to the degenerate probe mixture. All three of these clones were found to be false-positives that did not encode TNF-IP when the DNA sequence of the clones was determined.
- 28. This negative result was obtained despite the fact that one of the methods disclosed in the '953 patent was followed in performing the cloning and screening experiments.

- 29. Moreover, this negative result was obtained even though it was later demonstrated by others that U937 cells produce an mRNA encoding TNF-IP as part of a larger transcript (EP 422339; Appendix D).
- 30. These researchers also screened a randomly-primed colon cDNA library using a degenerate probe mixture containing the nucleotide base analog inosine (which was not described in the '953 patent) and obtained a positively-hybridizing longer cDNA fragment.
- 31. This longer cDNA sequence is designated C2 in the reference and is present in the Israeli Patent Application (Appendix C). The C2 sequence was incomplete and devoid of a translational start codon or a stop codon. The C2 sequence also includes a number of bases that are different from the claimed sequences (see sequence comparison, Appendix M). Thus, the C2 sequence appears to be a related but distinct sequence from the claimed sequence.
- 32. The C2 sequence was used as a second probe for screening the colon cDNA library. The '953 patent did not disclose production of a secondary probe comprising what appears to be a distinct sequence for screening a cDNA library.
- 33. Even when using the longer C2 sequence as a probe, the attempt to identify a full-length clone encoding TNF-IP failed in two experiments: the first being a rescreening of the colon library from which the C2 sequence had been obtained, and the second being the screening of an oligo dT-primed placental cDNA library.
- 34. It was not until the experiments set forth in the publication of Nophar et al. (Appendix E) were performed that these inventors were able to produce a TNF-IP encoding cDNA clone. However, this success was achieved well after the effective filing date of the '953 patent.
- 35. The experiments described in the Nophar *et al.* reference required screening a λ ZAP cDNA library constructed from randomly-primed cDNA produced from human CEM cells.
- 36. The CEM library used by Nophar et al. was not available at the effective filing date of the '953 invention (see the Clontech catalogue dated 1998; Appendix F). This library only became available after the effective filing date of the '953 patent (see the Clontech catalog dated 1989-1990; Appendix G).
 - 37. Nophar et al. does not teach why the CEM cDNA library was chosen, but CEM

cells apparently were <u>not</u> identified as being TNF-IP expressors using immunofluorescence or Western blotting, as described in the '953 patent.

- 38. The evidence from the Nophar *et al.* reference demonstrates that the specification of the '953 patent did not enable one having ordinary skill in the art to produce a recombinant DNA encoding TNF-IP.
- 39. Even the inventors of the '953 patent themselves were not able to isolate a TNF-IP encoding clone using the teachings of the '953 patent.

Attempts of Other Research Groups to Clone DNA

- 40. In addition, other well-funded, motivated researchers at companies such as Boehringer Ingelheim, Hoffman-La Roche, Genentech and Synergen attempted to clone the DNA. Although there were numerous failures by these groups to isolate a DNA molecule encoding TNF-IP using information at least as detailed as in the '953 patent, not one of those groups successfully isolated a DNA molecule encoding TNF-IP by simply using the information on the face of '953 patent.
- 41. As set forth below, each of these research groups was required to creatively resolve critical issues left unresolved by the teachings of the '953 patent. For example, each of the research groups (a) used affinity chromatography, a biochemical purification protocol different from that reported in the '953 patent; (b) generated and used additional amino acid sequence information to generate hybridization probes; and (c) identified and screened cell sources not identified in nor identifiable from the '953 patent.
- 42. For example, researchers at Hoffman-LaRoche reported their cloning of cDNA encoding the TNF receptor (Loetscher et al., 1990, Cell 61: 351-359; Appendix H).
- 43. These researchers reported that "conventional cloning" using "relatively short, fully degenerate or longer best-guess oligonucleotides" for screening oligo dT-primed cDNA libraries was "technically difficult" and had not resulted in cloning cDNA encoding the TNF receptor or TNF-IP (p.356). Sources of mRNA for preparing cDNA libraries screened in these failed experiments included human placental tissue and the human cell line HL60 (p. 357).
- 44. These researchers only successfully obtained a cDNA clone encoding the TNF receptor after producing a 78bp cDNA fragment using the polymerase chain reaction (PCR) with

an N-terminal sequence and an internal peptide fragment from an affinity-purified preparation, and screening a human placenta $\lambda gt11$ cDNA library with this probe.

- 45. The '953 patent does not disclose the sequence of any internal peptide fragment, and thus not the specific fragment used by the Hoffman-LaRoche researchers, for preparing oligonucleotide probes. Moreover, the '953 patent makes no mention of using PCR to produce either cDNA or probes for screening cDNA.
- The researchers did not state that the placental tissue was identified as being TNF-IP expressors using immunofluorescence or Western blotting, as described in the '953 patent.
- 47. The '953 patent does not identify placental tissue as providing a cell source of TNF-IP expressors.
- 48. Thus, the evidence from the Loetscher *et al.* reference is that these researchers were successful in obtaining a cDNA clone encoding TNF-IP only when (a) they obtained a 78bp, PCR-produced cDNA fragment, which was not disclosed in the patent and (b) they used the fragment to screen a human placenta λgt11 cDNA library, which was not disclosed in the '953 patent.
- 49. In another example, researchers at Boehringer Ingelheim screened oligo dT-primed cDNA libraries prepared from placenta and human cell lines U937, Hs913T and HeLa using mixtures of degenerate oligonucleotide probes derived from the N-terminal amino acid sequence from an affinity-purified preparation disclosed in the '953 patent (European Patent Application EP417563; Appendix I).
 - 50. These experiments failed to produce a cDNA encoding TNF-IP.
- 51. It was only when these researchers screened an Hs913T library with a PCR fragment using the amino acid sequence of an internal tryptic peptide fragment from an affinity-purified preparation that a cDNA encoding TNF-IP was obtained.
- 52. The '953 patent does not disclose the sequence of any internal peptide fragment, and thus not the specific fragment used by the Boehringer Ingelheim researchers, for preparing oligonucleotide probes. Moreover, the '953 patent does not disclose cDNA cloning or screening methods using the PCR.
- 53. The researchers did not state that the Hs913T cells were identified as being TNF-IP expressors using immunofluorescence or Western blotting, as described in the '953 patent.

- 54. The '953 patent does not identify Hs913T cells as being cells that express TNF-IP.
- 55. Thus, the evidence from European Patent Application EP417563 is that these researchers were successful in obtaining a cDNA clone encoding TNF-IP only when (a) they obtained a PCR-produced fragment corresponding to an internal, protease-generated peptide, which was not disclosed in the '953 patent and (b) they used the fragment to screen an oligo dT-primed Hs913T cDNA library, which was not disclosed in the '953 patent.
- 56. In yet another example, researchers at Genentech reported their cloning of cDNA encoding the TNF receptor (Schall et al., 1990, Cell 61: 361-370; Appendix J).
- 57. These researchers produced two purified, internal proteolytic fragments from an affinity-purified preparation that were used to prepare oligonucleotides for screening placental and HL60 cDNA libraries made in λgt10 (p. 368)
- 58. The Genentech researchers obtained four positively-hybridizing clones from a randomly-primed library and a single positively-hybridizing clone from an oligo dT-primed library.
- 59. The oligonucleotide probes used to screen these libraries were not degenerate, being specifically derived from less polymorphic regions of the protein and using the codon usage rules of Lathe *et al.* (p. 368).
- 60. The '953 patent does not identify the sequence of any internal peptide fragment for preparing non-degenerate oligonucleotide probes, and thus not the specific fragment used by the Genentech researchers, for preparing oligonucleotide probes
- 61. The researchers did not state that the HL60 cells were identified as being TNF-IP expressors using immunofluorescence or Western blotting, as described in the '953 patent.
 - 62. The '953 patent does not identify HL60 cells as being cells that express TNF-IP.
- 63. Thus, the evidence from the Schall et al. reference is that these researchers were successful in obtaining a cDNA clone encoding TNF-IP only when they (a) used a cDNA library produced from a tissue (placenta) and a cell line (HL60) that were not first identified as TNF-IP expressors using antibodies according to the teachings of the '953 patent and (b) a sequence encoding an internal peptide that was not identified in the patent.
 - 64. In a still further example, researchers at Synergen reported the cloning of cDNA

encoding the TNF receptor (EP 422339; Appendix D).

- 65. These researchers produced purified proteolytic fragments from an affinity-purified preparation that was used to prepare four degenerate oligonucleotide probes encoding N-terminal and internal peptide sequences for screening a genomic library (p. 23, ln. 32-52).
- 66. Initially an N-terminal probe identified eleven positive clones, with one clone hybridizing with all four degenerate probes (p. 23, ln. 55 through p. 24, ln. 14). The one clone was sequenced and encompassed significantly more of the N-terminal sequence (68 amino acids) from that which was taught in the '953 patent (p. 24, ln. 15-29).
- 67. The Synergen researchers used a probe encoding an internal peptide sequence of TNF-BP to screen an oligo dT-primed cDNA of U937 cells, from which three clones were identified (p. 24, ln. 49-53). The three plaques were confirmed with a probe encoding a second internal peptide sequence (p. 24, ln. 52-53).
- 68. The '953 patent does not disclose the sequence of any internal peptide fragment, and thus not the specific fragment used by the Synergen researchers, for preparing oligonucleotide probes.
- 69. The researchers did not identify U937 cells as being TNF-IP expressors using immunofluorescence or Western blotting, as described in the '953 patent.
- 70. The '953 patent does not identify U937 cells as being cells that express TNF-IP. In fact, a later-published reference by some of the inventors of the '953 patent support Applicants' position that the teachings of '953 may not have identified U937 cells as a suitable cell source.
- 71. The reference teaches anti-TNF-IP antibodies which were produced after immunization of mice with a TNF-IP preparation purified using a substantially different purification protocol that included affinity chromatography with immobilized TNF α (Engelmann et al., 1989, J. Biol. Chem. 264: 11970-11980; Appendix K). The monoclonal antibodies produced according to the Engelmann et al. reference failed to bind to U937 cells.
- 72. Thus, the evidence from the EP 422339 is that these researchers were successful in obtaining a cDNA clone encoding TNF-IP only when they used (a) a cDNA library produced from a cell line (HL60) that was not first identified as TNF-IP expressors using antibodies according to the teachings of the '953 patent and (b) sequence encoding an internal peptide that

was not identified in the patent.

Conclusion

- 73. In my opinion, the disclosure of the '953 patent would not have enabled one of ordinary skill in the art to obtain a DNA molecule encoding TNF-IP for at least the following reasons.
- 74. The '953 patent does not enable one of ordinary skill in the art to obtain a cDNA molecule encoding TNF-IP because, at the effective filing date of the patent, no tissue or cell line was known that expressed TNF-IP.
- 75. The fact that TNF-IP is a proteolytic cleavage product of the TNF receptor, and represents the extracellular domain of the receptor, was not known at the effective filing date of the '953 patent (see, for example, Wallach et al., 1989, Lymphokine Research 8: 361, which says:

"The cellular source of this urine-derived TNF binding protein remains to be elucidated." (Appendix L)

- 76. The protein prepared from the biochemical separation protocol described in the '953 patent is very impure, being >50% contaminated with other proteins.
- 77. This level of impurity would make the protein preparation described in the '953 patent difficult to use in preparing and screening antibodies specific for TNF-IP.
- 78. Also, this level of impurity would have led one of ordinary skill to believe that the peptide sequence disclosed in the '953 patent could not be relied upon as being related to the TNF-IP, since there was a better than even chance that this sequence was completely unrelated to the TNF inhibiting activity.
- 79. In addition, until a cDNA clone is isolated and a recombinant TNF-IP protein produced and characterized, it could not be established that the protein from which the N-terminal amino acid sequence was obtained in fact binds to TNF.
- 80. No cDNA libraries or sequence of probes encoding the putative N-terminal amino acid sequence were disclosed in the '953 patent.
- 81. The '953 patent only provides a limited amount of sequence information and an encyclopedia of experimental options. After having spent considerable time and effort in discovering that essential information was lacking in the patent, I would have been left with myriad potential research avenues but no specific guideposts of how to successfully navigate

them. In fact, each research group that successfully cloned the DNA relied on insights and methods that were not described in the '953 patent.

- 82. In summary, it is clear that the amino acid sequence disclosed in the patent and the absence of an identified cell source in the patent provided insufficient information to clone the DNA.
- 83. Thus, it is my opinion that one of ordinary skill in the art would not have been able to obtain a DNA molecule encoding TNF-IP using the methods set forth in the '953 patent.

I hereby declare further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Signed:	and most	
	John D. Mountz, M.D. Ph.D.	

Dated: 2/25/99

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE (Attorney Docket No. 98,385-A)

In the	Application of:)	
Hauptı	nann et al.) Examiner: G. Draper	
Serial I	No: 08/484,312) Group Art Unit: 1646	
Filed:	June 7, 1995)	66 HO31
For:	TNF Receptors, DNAs Coding for	TNF Binding Proteins and or them)))	H CENTS
	Commissioner for ngton, DC 20231	[ITTAL LETTER	AH III
	In regard to the	above-identified patent app	olication:	000
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		No additional fee is require Attached are two checks in	d. the amount of	99 MAR - 1 PM 12: 53
3.	Please charge any additional fees or credit over-payments to the Deposit Account No.13-2490.			
4.	<u>x</u> CERTIFICATE UNDER 37 CFR 1.18: The undersigned hereby certifies that this Transmittal Letter and this paper, as described in paragraph 1 hereinabove, are being hand delivered to: Examiner G. Draper in Group Art Unit 1646, Washington, D.C. 20231, on this 1st day of March 1999.			
Dated:	March 1, 1999	By:	John J. McDonnell Reg. 26,949	

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Appendix A

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JOHN MOUNTZ UAB

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Updated 08/20/98

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Consultant Scientist; Wyeth-Ayerst, Princeton, NJ

Consultant Scientist; LXR Biotechnologies, Inc., Richmond, CA

Consultant Scientist; Amgen Boulder, Inc., Boulder, CO

EDUCATION:

Institution	Degree	Year
Wright State University, Dayton, OH Michigan State University, East Lansing, MI Michigan State University, East Lansing, MI Ohio State University, Columbus, OH	BS MS PhD MD	1971 1971 1974 1978
•		1710

MILITARY SERVICE: N/A

LICENSURE:

Physician, North Carolina #23624 Physician, Alabama #13416

BOARD CERTIFICATION:

American Board of Internal Medicine - 1982

POSTDOCTORAL TRAINING:

Year	Degree	Institution
1974-1975	Postdoctoral Fellow	National Science Foundation East Lansing, MI
	Medical School Scholarships	William Craig-Orr and Mary Black-Orr Scholarship in Medicine
		Kenneth Wiseman Scholarship in Medicine
1978-1981	Intern and Resident	Internal Medicine Residency Program; North Carolina Baptist Hospital
1981-1982	Rheumatology Fellow	Bowman Gray School of Medicine
1982-1987	Medical Staff Fellowship	National Institutes of Health; National Institute of Arthritis, Diabetes, and Digestive and Kidney Diseases
1988-present	Director	Transgenic Mouse Facility, Birmingham Veterans Administration Medical Center

ACADEMIC APPOINTMENTS:

JOHN MOUNTZ UAB

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Year	Rank/Title	Institution
1985-1987	Research Assistant Professor of Medicine	Uniformed Services University of Health Sciences
1986-1987	Guest Researcher	National Institutes of Health; National Institute of Arthritis and
1987-1991	Assistant Professor of Medicine Division of Clinical Immunology and Rheumatology	Musculoskeletal and Skin Diseases UAB
1991-1994	Associate Professor of Medicine Division of Clinical Immunology and Rheumatology	UAB
1991-present	Associate Professor, Division of Gerontology and Geriatric Medicine	UAB
1991-present	Associate Professor, Cellular & Molecular Biology, Department of Microbiology (Joint appointment renewed for 3 years March, 1996)	UAB
1987-present	Staff Physician	Birmingham VAMC
1993-present	Scientist, Medical Center Joint Departments, Center for Aging	UAB
1994-present	Professor of Medicine, Division of Clinical Immunology and Rheumatology	UAB

HONORS AND AWARDS:

B.S., Summa Cum Laude
M.S., Summa Cum Laude
Phi Eta Tau - Scholastic Honor Society
Elected to Who's Who of Birmingham - 1990

AWARDS FOR RESEARCH ACCOMPLISHMENTS:

Senior Rheumatology Scholar Award - 1986 Howard and Martha Holley Research Prize in Rheumatology - March 13, 1992

MEMBERSHIPS AND OFFICES IN PROFESSIONAL SOCIETIES:

American College of Physicians - 1978
American Association of Immunology - 1985
American College of Rheumatology - 1986
Southern Society of Clinical Investigation - 1988
American Federation for Clinical Research - 1990

American Society of Internal Medicine - 1990 American Society of Clinical Investigation - 1992 Southern Medical Association - 1992 Association of American Physicians - 1997

COUNCILS AND COMMITTEES:

COMMITTEES

Anhritis and Prostaglandins Research Challenge

Advisory Meeting. Searle Pharmaceuticals, Chicago, IL. May 3-6, 1990.

Veterans Administration Transgenic Mouse Facility Scientific Advisory Committee (1990-)

Veterans Administration Medical Center Transgenic Mouse Subcommittee (1990-)

Veterans Administration Animal Use Committee (1991-)

Department of Medicine Research Committee (1991-)

Admissions Committee, School of Medicine, University of Alabama at Birmingham,

Department of Medicine, MD, PhD. Training Program (1991-)

Research Council of American College of Physicians (1993-1996)

Geriatrics Research Center Advisory Committee (1993-)

Chairman, Adhesion Molecule and Cell Matrix Center Director Search Committee (1993)

Committee for Revision of American College of Rheumatology Patient Care Summary (1993-1994)

Research Committee of the ACR Research and Education Foundation (1994-)

Co-Chairs, Task Force, Committee, National Institute on Aging, National Institute of Allergy and Infectious Diseases Task Force on Immunology and Aging

Veterans Administration Merit and Career Development Review Committee; Immunology Study Section, (1995-1998)

Chairman, Department of Medicine Research Committee (1996-)

UAB/Sankyo Co., Ltd., Program for Rheumatic Diseases Oversight Committee (1996)

UNIVERSITY ACTIVITIES:

Department of Medicine Research Committee (1991-)

Veterans Administration Transgenic Mouse Facility Scientific Advisory Committee (1990-)

Veterans Administration Medical Center Transgenic Mouse Subcommittee (1990-)

Veterans Administration Animal Use Committee (1991-)

Admissions Committee, School of Medicine, University of Alabama at Birmingham,

Department of Medicine, MD, PhD. Training Program (1991-)

Geriatrics Research Center Advisory Committee (1993-)

Chairman, Adhesion Molecule and Cell Matrix Center Director Search Committee (1993)

Chairman, Department of Medicine Research Committee (1996-)

UAB/Sankyo Co., Ltd., Program for Rheumatic Diseases Oversight Committee (1996)

02/12/99

JOHN MOUNTZ UAB

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EDITORIAL BOARDS/AD HOC REVIEWER:

Editorial Board, Arthritis & Rheumatism; 1988 - 1993 Editorial Board, Cellular Immunology; 1996 - 2000 Reviewer, American Association of Immunologist

Reviewer, American Journal of Medicine

Reviewer, Autoimmunity

Reviewer, Blood

Reviewer, Antiviral Research

Reviewer, Clinical Chemistry

Reviewer, Gastroenterology

Reviewer, Human Immunology

Reviewer, International Immunology

Reviewer, Immunity

Reviewer, Immunology Today

Reviewer, Journal of Clinical Immunology

Reviewer, Journal of Clinical Investigation

Reviewer, Journal of Rheumatology

Reviewer, Science

Reviewer, Transgenes and Transgenic Mice

Reviewer, Transgenic Technology

Reviewer, FASEB Journal

Reviewer, Veterans Administration Health Services and Research Administration, Career Development Program Applications, Study Section, National Arthritis Foundation (1991-1993)

Immunological Sciences Study Section Ad Hoc Reviewer, NIAID, June 9-11, 1993 External Reviewer, Medical Research Service, Department of Veterans Affairs, 1995-1997 Ad Hoc Reviewer, The Israel Science Foundation, Israel Academy of Science and Humanities, Jerusalem, Israel, March, 1996. Grant Reviewer, The Arthritis and Rheumatism Council, 1997 Reviewer, The Wellcome Foundation, 1997 Immunological Sciences Study Section, 1997-2000

Site Visits:

Special Project Reviewer for NIH - U. California at San Diego Multipurpose and Musculoskeletal Disease Center (MAMDC) 1991 Special Project Reviewer for NIH, U. Texas at Dallas, MAMDC. 1991 Special Project Reviewer, Boston University School of Medicine, March 1996 Reviewer, Cornell Aging Program Project, December, 1996 Reviewer, University of Michigan Aging Program Project, February 1998. Reviewer, "Cellular Senescence and Control of Proliferation" Allegheny University of the Health Sciences, Phildelphia, PA, July 28, 1998.

MAJOR LECTURES AND VISITING PROFESSORSHIPS:

- 1. "Therapies which reduce lymphadenopathy and autoimmunity of MRL/lpr Mice: Implications for the Pathogenesis and Treatment of RA and SLE," New York Academy of Sciences, November 12, 1987.
- 2. "The Role of T Cells in Autoimmune Disease," Lowe Southeastern Conference on Rheumatic Diseases, September 28, 1990.
- 3. "Update on Arthritis Research." Arthritis Foundation Dinner, 3rd Humanitarian Award Ceremony, September 21, 1990.
- 4. "A Superantigen Model for Arthritis in V,8 TCR Transgenic *lpr/lpr* Mice," Keystone Symposia on the Molecular Biology and the Immunopathogenesis of Rheumatoid Arthritis, March 16, 1991.
- 5. "Loss of T-cell Anergy in Autoimmunity Mice," Bar Harbor Symposia on MHC and TCR in Autoimmune Disease, October 18-20, 1991, Bar Harbor, Maine.
- 6. "T Cell Tolerance Defects in Transgenic Mice," Holley Symposia at Regional ACR Meeting, March 13, 1992.
- 7. "ETn Mutation in Fus CD2-fas Correction of lpr Mice," Keystone Symposia on the Molecular Biology and the Immunopathogenesis of Rheumatoid Arthritis, March 16, 1993.
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Appendix B

IS005695953A

United States Patent 1191

Wallach et al.

[11] Patent Number:

5,695,953

[45] Date of Patent:

Dec. 9, 1997

[54] DNA THAT ENCODES A TUMOR NECROSIS FACTOR INHIBITORY PROTEIN AND A RECOMBINANT METHOD OF PRODUCTION

[75] Inventors: David Wallach. Rebovot, Israel;

Hartmut Engelmann, Munich, Germany; Dan Aderka, Holon;

Menachem Rubinstein, Givat Schmuel,

both of Israel

[73] Assignee: Yeda Research and Development Co.

Ltd., Rebovot, Israel

[21] Appl. No.: 876,828

[22] Filed: Apr. 30, 1992

Related U.S. Application Data

[63] Continuation of Ser. No. 243,092, Sep. 12, 1988, abandoned.

[30] Foreign Application Priority Data

15: 536/23.5

[56]

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Primary Examiner-Garnette D. Draper Attorney, Agent, or Firm-Browdy and Neimark

[7] ABSTRACT

Tumor Necrosis Factor (TNF) Inhibitory Protein is isolated and substantially purified and the DNA that encodes the TNF inhibitory protein, vectors, host cells, and a recombinant method for producing the encoded protein are also set forth. It has the ability to inhibit: (a) the binding of TNF to its receptors, and (b) the cytotoxic effect of TNF. TNF Inhibitory Protein and salts, functional derivatives and active fractions thereof can be used to antagonize the deleterious effects of TNF.

15 Claims, 5 Drawing Sheets

Appendix C

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
- (i) APPLICANT: Hauptman, et al.
- (ii) TITLE OF INVENTION: THE RECEPTORS, THE BINDING BINDING PROTEINS, AND DNAs CODING FOR THEM
- (iii) NUMBER OF SEQUENCES:64
- (iv) CORRESPONDENCE ADDRESS:
- (A) ADDRESSEE: John J. McDonnell
- (B) STREET: 300 S. Wacker Drive
- (C) CITY: Chicago
- (D) STATE: IL
- (E) COUNTRY: USA
- (F) ZIP:60606
- (v) COMPUTER READABLE FORM:
- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- (D) SOFTWARE: ASCII
- (vi) CURRENT APPLICATION DATA:
- (A) APPLICATION NUMBER: 08/484,312
- (B) FILING DATE: June 7, 1995
- (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
- (A) NAME: John J. McDonnell
- (B) REGISTRATION NUMBER: 26,949
- (C) REFERENCE/DOCKET NUMBER: 98,385-A
- (ix) TELECOMMUNICATION INFORMATION:
- (A) TELEPHONE: 312-913-0001
- (B) TELEFAX:312-913-9808
- (2) INFORMATION FOR SEQ ID NO:1:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 1365 bases
- (B) TYPE:nucleic acid
- (C) STRANDEDNESS:single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GCTGTTGGTG GGAATATACC CCTCAGGGGT TATTGGACTG GTCCCTCACC 100 TAGGGGACAG GGAGAAGAGA GATAGTGTGT GTCCCCAAGG AAAATATATC 150 CACCCTCAAA ATAATTCGAT TTGCTGTACC AAGTGCCACA AAGGAACCTA 200 CTTGTACAAT GACTGCCAG GCCCGGGGCA GGATACGGAC TGCAGGGAGT 250 GTGAGAGCGG CTCCTTCACC GCTTCAGAAA ACCACCTCAG ACACTGCCTC 300 AGCTGCTCCA AATGCCGAAA GGAAATGGGT CAGGTGGAGA TCTCTCTTG 350 CACAGTGGAC CGGGACACCG TGTGTGGCTG CAGGAAGAAC CAGTACCGGC 400 ATTATTGGAG TGAAAACCTT TCCAGTGCT TCAATTGCAG CCTCTGCCTC 450 AATGGGACCG TGCACCAGAGA AACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTCTTC TAAGAGAAAA CGAGTGTCT TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACGAGAG AAACAGAACA CCGTGTGCAC 500 AATGTTAAGG GCACTGAGG TGCACGAGA TGTGCCTC CCAGGAGAAAA CGAGTGCTC TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACGAGAG TGTGCCTCC CCAGATTGAG 600 AATGTTAAGG GCACTGAGG TCCAAGGACC ACAGTGCTG TGCCCCTGGT 650 AATGTTAAGG GCACTGAGG TCCAAGGACC ACAGTGCTG TGCCCCTGGT 650 AATGTTAAGG GCACTGAGG TCCAAGCTCT ACTCCATTG TGCCCCTGGT 650 AATGTTAAGA AAGCCTGAG TCCAAGCTCT ACTCCATTG TTGTGGGAAA 750 ACGGCCCAACCC CCAAGCTTCA GTCCCACCTG CCCAGATTGAG 660 AATGTTACTT TCCAGTGCA GAGCCCC TCCAAGCCCC 660 CCCAAGCTCC CCAAGCTTCA CCCACCTC CCCACCTGG 750 CCCAGGTGCC CCCAAGCTT TCCACCTTCA CCTCCAGCCC CACCCTGG 850 CCCCAACCC CCTTCAGAAG TCCAACCTTCA CCTCCAGCTC CACCCCTG 680 CCCAGGTGACT TCCACCTTCA CCTCCAGCCC CACCCTGG 750 CCCAGGTGAC TCCACCCACT TCCACCTCC CCCAGAGAG TGGCACCAC 950 CCCAGGTGAC TCCACCCACT TCCACCTCC CCCAGAGAG TGGCACCAC 1000 CCCAGTGCC TGGAAGAA TCCTTGCGAC AGCCCCAAA GCCACAGAG 1050 CCCAGTGCC TGGAAGAAA TCCTTGCGAC AGCCCCAAA GCCACAGAG 1150 CCCAGTGCC TGGAAGAGA TCCTTGCGAC AGCCCCAAA GCCACAGAG 1150 CCCAGTGCC TGGAAGAG TCCAGCCC AGGCGCG CCAGAGAC CCCAAGAC 1150 CCCAGTGCC TGGAAGAG TCCACCC TCCGACCCC TCCGACCCC 1100 CCCAGTGCC TGGAAGAG TCCACCC TCCGCGC CCGGAGAGAC CCCACAGAC 1250 CCCAGGAGAC TCGAGAGAC TCCCCCCCCCC CCGAGAGACC CCCACACACCC TCGCCC TCGGCCCC TCGGCCCC TCGGCCCC TCGGCCCC TCGGCCCCCCCC	ATGGGCCTCT	CCACCGTGCC	TGACCTGCTG	CTGCCACTGG	TGCTCCTGGA	50
CACCCTCAAAATAATTCGATTTGCTGTACCAAGGGCCACAAAGGAACCTA200CTTGTACAATGACTGTCCAGGCCCGGGGCAGGATACGGACTGCAGGGAGT250GTGAGAGAGCGCTCCTTCACCGCTTCAGAAAACACCTCAGACACTGCCTC300AGCTGCTCCAAATGCCGAAAGGAAATGGGTCAGGTGGAGATCTCTTCTTG350CACAGTGGACCGGGACACCGTGTGTGGCTGCAGGAAGAACCAGTACCGGC400ATTATTGGAGTGCACCTCTCCTGCCAGGAGAAACAGAACACCGTGTGCAC500CTGCCATGCAGGTTTCTTCTAAGAGAAAACGAGTTGTGCTCCTGTAGTA550ACTGTAAGAAAAGCCTGGAGTGCACGAGAGTGTGCCTACCCCAGATTGAG600AATGTTAAGGGCACTGAGGACTCAGGCACACAGTGCTGTGCCCTGGT650CATTTTCTTTGGTCTTTGCCTTTTATCCCTCCTCTTCATTGGTTTAATGT700ATCGCTACCAACGGTGGAAGTCCAAGCTCTACTCCATTCACTAGGCACC800GCCCCAAACCCAAGCTTCAAGCCTTCACCACACCTGG800GCTTCAGTCCCGTGCCAGTTCCCACCTCCAGCCTTCACCCACCCTTGC950CCTGTGGACTGTCCCAACTTTGCGCCTCCCGCAGAGAGATGCCACCCT950CCTGCGTGACTGTCCCAACTTTGCGGCTCCCGCAGAGAGGTGCCACCCA950CCTGTGGGCGTGACCACCTCCTTGCGACAGCCCTCACATCCCACCTCAGCCCTCACA100CCCGTTGCGCCCTTCAGAGATCCTTGCGACAGCCCTCACAAGCACCACA110CCCGTTGCGCCCTGAGGCGCCCTAGGCCCAGCACCACA1	GCTGTTGGTG	GGAATATACC	CCTCAGGGGT	TATTGGACTG	GTCCCTCACC	100
CTTGTACAATGACTGTCCAGGCCCGGGGCAGGATACGGACTGCAGGGAGT250GTGAGAGCGGCTCCTTCACCGCTTCAGAAAACCACCTCAGACACTGCCTC300AGCTGCTCCAAATGCCGAAAGGAAATGGGTCAGGTGGAGATCTCTTCTTG350CACAGTGGACCGGGACACCGTGTGTGGCTGCAGGAAGAACCAGTACCGGC400ATTATTGGAGTGAAAACCTTTTCCAGTGCTTCAATTGCAGCCTCTGCCTC450AATGGGACCGTGCACCTCTCCTGCCAGGAGAAACAGAACACCGTGTGCAC500CTGCCATGCAGGTTTCTTTCTAAGAGAAAACGAGTGTGTCTCCTGTAGTA550ACTGTAAGAAAAGCCTGGAGTGCACGAAGTTGTGCCTACCCCAGATTGAG600AATGTTAAGGGCACTGAGGACTCAGGCACCACAGTGCTGTTGCCCCTGGT650CATTTTCTTTGGTCTTTGCCTTTTATCCCTCCTCTTCATTGGTTTAATGT700ATCGCTACCAACGGTGGAAGTCCAAGCTCTACTCCATTGTTTGTGGGAAA750TCGACACCTGAAAAAGAGGGGGAGCTTGAAGGAACTACTACTAAGCCCCT800GCTTCAGTCCCCAAGCTTCAGTCCCACTCCAGGCTTCACCCCCACCCTGG850GCTTCAGTCCGTGCCCAACTTGCACCTTCACCTCCAGCTCCACCTATACC900CCCGGTGACCTCCTTGCGACAGCCCCCACAAGCCACAGAGC1000TCCCCAACCCCCTTCAGAAGTCCTTGCGACAGCACACAGAG1000CCCGTTGCGCCCTTCAGAAGTCCTGCGCGCCTAGGGCTGAGCACAGAGC1100CCCGTTGCGCTGGAAGGACTCCGGCGCGCCCGAGGCCAC1250 <tr< td=""><td>TAGGGGACAG</td><td>GGAGAAGAGA</td><td>GATAGTGTGT</td><td>GTCCCCAAGG</td><td>AAAATATATC</td><td>150</td></tr<>	TAGGGGACAG	GGAGAAGAGA	GATAGTGTGT	GTCCCCAAGG	AAAATATATC	150
GTGAGAGCGG CTCCTTCACC GCTTCAGAAA ACCACCTCAG ACACTGCCTC 300 AGCTGCTCCA AATGCCGAAA GGAAATGGGT CAGGTGGAGA TCTCTTCTTG 350 CACAGTGGAC CGGGACACCG TGTGTGGCTG CAGGAAGAAC CAGTACCGGC 400 ATTATTGGAG TGAAAACCTT TTCCAGTGCT TCAATTGCAG CCTCTGCCTC 450 AATGGGACCG TGCACCTCC CTGCCAGGAG AAACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTTCTTC TAAGAGAAAA CGAGTGTGTC TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACCGAGAG AAACAGAACA CCGTGTGCAC 500 AATGTTAAGG GCACTGAGG TGCACCGAGAGT TGTGCCTACC CCAGATTGAG 600 AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 CGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCAC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGCC CGCAGAGAGG TGGCACCAC 950 CTAACACCCC CCTTCAGAAG TGGGAGGACA GCCCCCACAC GCCCCCAACC CCTAGACCCC ACCCTTGCG CCCCCACACC CCTAGACCCC ACCCTTGCG CCCCCCCACACC CCCCCCCCC GACCCCCC GACCCCCCC CCCCCCCC	CACCCTCAAA	ATAATTCGAT	TTGCTGTACC	AAGTGCCACA	AAGGAACCTA	200
AGCTGCTCCA AATGCCGAAA GGAAATGGGT CAGGTGGAGA TCTCTTCTTG 350 CACAGTGGAC CGGGACACCG TGTGTGGCTG CAGGAGAAC CAGTACCGGC 450 ATTATTGGAG TGAAAACCTT TTCCAGTGCT TCAATTGCAG CCTCTGCCTC 450 AATGGGACCG TGCACCTCTC CTGCCAGGAG AAACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTTCTTTC TAAGAGAAAA CGAGTGTGT TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACGAGA TGTGCCTACC CCAGATTGAG 600 AATGTAAGA AAGCCTGGAG TGCACGAAGT TGTGCCTACC CCAGATTGAG 650 AATGTTAAGA AAGCCTGGAG TCCAGGACC ACAGTGCTGT TGCCCTGGT 650 AATGTTAAGA AAGCCTGGAG CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 700 AATGTTAATGT 700 ACTGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTG TTGTGGGAAA 750 ACGGCCCAACA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 ACGCCCAACC CCAAGCTCA AAAAAGAGAG GGAACTACTA CTAAGCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACCTC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTC CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACCT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACCT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGG GCTGACCCA TCCTTGCGAC AGCCCTCGC TCCGACCCCA 1000 CCCGGTGACC CCTTCAGAAG TGGGAGGACA GCCACAGAGC 1050 CTAGACACCT ATGACCCCG GACGCTGTAC GCCCCACAA GCCACAGAGC 1050 CTAGACACCT ATGACCCCG GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGCG CCTAGGGCTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGCG CCTAGGGCTG AGAACGTGCC 1100 CCCGTTGCG TGGAAGGAAT TCGTGCGCG CCTAGGGCTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGCG CCTAGGGCTG AGAACGTGC 1100 CCCGTTGCGC TGGGAAGGAT TCGTGCGCG CCTAGGGCTG AGAACGTGC 1100 CCCGTTGCGC TGGGAAGGAT TCGTGCGGC CCTAGGGCTG CCGAGGCCAC 1250 GCTGGAGCCAC TGGGAGCAC TGGGAGCAC TGGGAGCAC TGGGAGGCAC TCGGAGGCCAC TCGGAGGCCAC TCGGAGGCCAC TCGGAGGCCAC TCGGAGGCCAC TCGGAGGCCAC TCGGAGGCAC TCGGAGGCAC TCGGAGGGCAC TCGGAGGGACAC CCGGCGGCAC TCGGAGGGCAC TCGGAGGGACAC CCGGCGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGACAC CCGGCGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGCAC TCGGAGGGACAC TCGGAGGGCAC TCGGAGGGACAC	CTTGTACAAT	GACTGTCCAG	GCCCGGGGCA	GGATACGGAC	TGCAGGGAGT	250
CACAGTGGAC CGGGACACCG TGTGTGGCTG CAGGAAGAC CAGTACCGGC ATTATTGGAG TGAAAACCTT TTCCAGTGCT TCAATTGCAG CCTCTGCCTC AATGGGACCG TGCACCTCTC CTGCCAGGAG AAACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTTCTTTC TAAGAGAAAA CGAGTGTGT TCCTGTAGTA ACTGTAAGAA AAGCCTGGAG TGCACGACCT TGCACGACC ACAGTGTGT CCCAGATTGAG AACGTAGCA CCCAGATTGAG AACTGTAAGAA AAGCCTGGAG TGCACGAAGT TGTGCCTACC CCAGATTGAG AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCACCCA ACAGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 GGCCCAAAC CCAAGCTTCA GGCCCTAGC AAAAAGAGGG GGACCTTGAA GGAACTACTA CTAAGCCCCT 850 GCTTCAGTCC CGTGGCCCAGT TCCACCTTCA CCCCACCCTGG 850 CTATCAGGC CCCACCCTGG CCCACCCTGG CCCCACCCTGG CCCCACCCTGG CCCCACCCC CCCACCCTGG CCCCACCCC CCCACCCTGC CCCCACCCC CCCACCCTGC CCCCACCCC CCCACCCCC CCCACCCCC CCCACCCCC CCCCCC	GTGAGAGCGG	CTCCTTCACC	GCTTCAGAAA	ACCACCTCAG	ACACTGCCTC	300
ATTATTGGAG TGAAAACCTT TTCCAGTGCT TCAATTGCAG CCTCTGCCTC 450 AATGGGACCG TGCACCTCTC CTGCCAGGAG AAACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTTCTTTC TAAGAGAAAA CGAGTGTGC TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACCGAGGAG TGTGCCTC CCAGATTGAG 660 AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCCCCCCACAA GCCACAGAGC 1050 CTAGACACTC ATGACCCCG GACGCTGTAC GCCGCGCGC AGAACGTGC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGC CCTAGGGCTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCACCACG 1150 AGATCGATCG TGGCACCTG GAGACGGC CCTAGGGCTG CGAGGCCCA 1200 TACAGCATGC TGGCACCTG GAGACGGC CCTAGGGCTG CGAGGCCCA 1200 TACAGCATGC TGGCGACCTG GAGACCGC CCTAGGGCTG CCGAGGCCCA 1200 TACAGCATGC TGGCGACCTG CAGAACGGC CCTAGGGCTG CGAGGCCCA 1200 TACAGCATGC TGGCGACCTG CAGAACGGC CCTAGGGCTG CGAGGCCCA 1200 TACAGCATGC CTGGGACCTG CAGAACGGC CCTAGGGCTG CCGAGGCCCA 1250 GCTGGAGCTG CTGGGACCGC CCGCCCCC CCGCCCCC CCGCCCCCC 1300	AGCTGCTCCA	AATGCCGAAA	GGAAATGGGT	CAGGTGGAGA	TCTCTTCTTG	350
AATGGGACCG TGCACCTCTC CTGCCAGGAG AAACAGAACA CCGTGTGCAC 500 CTGCCATGCA GGTTTCTTTC TAAGAGAAAA CGAGTGTGTC TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACCGAAGT TGTGCCTACC CCAGATTGAG 600 AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACAGTGCTGT ACTCCATTGT TTGTGGGAAA 750 TCGACACCT AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTC CGTGCCCAACC CGAGCTTCA CCCACCTCC CGCAGCACC CACCTATACC 900 CCCGGTGACT GTCCCAACCT TGCGGCACC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCC 950 CTAGACCCC CCTTCAGAAG TGGGAGGACA GCCCCCACAA GCCACAGAGC 1000 CCCGTTGCG ATGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGAACGTGCC 1150 AGATCGATCG TGGCACCTG GAGACCGCG CCTAGGGCTG CGAGGCCCAA 1200 TACAGCATGC TGGCACCTG GAGACCGCG CCTAGGGCTG CGAGGCCCA 1250 GCTGGAGCTG CTGGGACCCC TGCCGCCC CGCAGAGCCC 1250 GCTGGAGCTG CTGGGACCAC TGCTGGAGCCC CTGGGCCCC CCGCGCCCC CTGGGCCCC CTGGGCCCC TGGGCCCC TGGGCCCC TGGGAGCCAC 1250 GCTGGAGCCT CTGGGACCCC TGCCCCCCC CCGCCCCC CCGCCCCC TGGGCCCC TGGGCCCC CTGGGAGCCAC 1250 GCTGGAGCCAC CTGGAGCCCC CCGCCCCCC CCGCCCCCC CCGCCCCCC CCGCCCCCC	CACAGTGGAC	CGGGACACCG	TGTGTGGCTG	CAGGAAGAAC	CAGTACCGGC	400
CTGCCATGCA GGTTTCTTC TAAGAGAAAA CGAGTGTGTC TCCTGTAGTA 550 ACTGTAAGAA AAGCCTGGAG TGCACGAAGT TGTGCCTACC CCAGATTGAG 600 AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGCG CCTAGGGCTG AGCACCACG 1150 AGATCGATCG CTGGAGCCTG CAGAACGGC GCTGCCTGC CGAGGCCCAC 1250 GCTGGAGCTG CTGGGACCCC CAGAACGGC CCTAGGGCTG CCGAGGCCCAC 1250 GCTGGAGCTG CTGGGACCCC CAGAACGGC CCTGGCC CTGGGCCCC 1300 TACAGCATGC CTGGGACCCG CAGAACGGC CCTGCCCCCCCCC CTGGGCCCC 1300 TGGAGGACAT CGAGGAGCC CTTTGCCGCC CCCCCCCCC CCCCCCCCC 1300	ATTATTGGAG	TGAAAACCTT	TTCCAGTGCT	TCAATTGCAG	CCTCTGCCTC	450
ACTGTAAGAA AAGCCTGGAG TGCACGAAGT TGTGCCTACC CCAGATTGAG 650 AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 650 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCG GACGCTGTAC GCCGTGGTG AGAACGTGC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCACCACG 1150 AGATCGATCG GCTGGACCTG CAGAACGGCC GCTGCCTGCG CGAGGCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGCG ACGCCGCGC GCGAGGCCAC 1250 GCTGGAGCACT TGGGGACGC TGCTCCGCG CTGCGCCCCCC 1300 TGGAAGGACAT CCTGGGACCC CCGCCCCCC CCGCCCCCC 1300	AATGGGACCG	TGCACCTCTC	CTGCCAGGAG	AAACAGAACA	CCGTGTGCAC	500
AATGTTAAGG GCACTGAGGA CTCAGGCACC ACAGTGCTGT TGCCCCTGGT 700 CATTTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCG GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGC CCTAGGGCTG AGCACCACG 1150 AGATCGATCG TGGAAGGAAT TCGTGCGGC CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG TGGCGACCTG GAGGCGGC ACGCCGCGC CGAGGCCAC 1250 GCTGGAGCTG CTGGGACCCC CCGCGCC CCGCGCC CTGGGCTGC 1300 TGGAAGGACAT CGAGGAGGC CCTTGCGCC CCGCCCCC 1300	CTGCCATGCA	GGTTTCTTTC	TAAGAGAAAA	CGAGTGTGTC	TCCTGTAGTA	550
CATTTCTTT GGTCTTTGCC TTTTATCCCT CCTCTTCATT GGTTTAATGT 700 ATCGCTACCA ACGGTGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCC CGAGGCCCA 1200 TACAGCATGC TGGCGACCTG GAGGCCGCC ACGCCGCGC GCGAGGCCCA 1200 TACAGCATGC TGGCGACCTG GAGGCCGCC CCTGCCC CGAGGCCCAC 1250 GCTGGAGCTG CTGGGACCCC CCGCCGCC CCGCCCCC 1300 TGGAGGACAT CGAGGAGGC CTTTGCCGCC CCGCCCCC CCCCCCCC 1300	ACTGTAAGAA	AAGCCTGGAG	TGCACGAAGT	TGTGCCTACC	CCAGATTGAG	600
ATCGCTACCA ACGGTGGAAG TCCAAGCTCT ACTCCATTGT TTGTGGGAAA 750 TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCC CGAGGCCCA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACCCC CCTTTGCCGCC CCGCCCCCC CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCCCCC CCCCCCCCC 1350	AATGTTAAGG	GCACTGAGGA	CTCAGGCACC	ACAGTGCTGT	TGCCCCTGGT	650
TCGACACCTG AAAAAGAGGG GGAGCTTGAA GGAACTACTA CTAAGCCCCT 800 GGCCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCGCCC CCCGCCGCG 1350	CATTTTCTTT	GGTCTTTGCC	TTTTATCCCT	CCTCTTCATT	GGTTTAATGT	700
GGCCCAAAC CCAAGCTTCA GTCCCACTCC AGGCTTCACC CCCACCCTGG 850 GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACTT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCACA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACCCC CCTTGCGCG CCGCGCCCCCCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCCCCC 1350	ATCGCTACCA	ACGGTGGAAG	TCCAAGCTCT	ACTCCATTGT	TTGTGGGAAA	750
GCTTCAGTCC CGTGCCCAGT TCCACCTTCA CCTCCAGCTC CACCTATACC 900 CCCGGTGACT GTCCCAACCT TGCGGCTCC CGCAGAGAGG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCACAA GCCACAGAGC 1050 CTAGACACCG ATGACCCCG GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACCCG TGCTCCGCG CCGCGCCCC CCGCCGCG 1350	TCGACACCTG	AAAAAGAGGG	GGAGCTTGAA	GGAACTACTA	CTAAGCCCCT	800
CCCGGTGACT GTCCCAACTT TGCGGCTCCC CGCAGAGAG TGGCACCACC 950 CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCGCCC CCCGCCGCG 1350	GGCCCCAAAC	CCAAGCTTCA	GTCCCACTCC	AGGCTTCACC	CCCACCCTGG	850
CTATCAGGGG GCTGACCCCA TCCTTGCGAC AGCCCTCGCC TCCGACCCCA 1000 TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCGCCC CCCGCCGCG 1350	GCTTCAGTCC	CGTGCCCAGT	TCCACCTTCA	CCTCCAGCTC	CACCTATACC	900
TCCCCAACCC CCTTCAGAAG TGGGAGGACA GCGCCCACAA GCCACAGAGC 1050 CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCGCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACCGC TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCGCCC CCCGCCGCG 1350	CCCGGTGACT	GTCCCAACTT	TGCGGCTCCC	CGCAGAGAGG	TGGCACCACC	950
CTAGACACTG ATGACCCCGC GACGCTGTAC GCCGTGGTGG AGAACGTGCC 1100 CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCCCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCG ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCCGCC CCGCCCCCC CCCCCCGCG 1350	CTATCAGGGG	GCTGACCCCA	TCCTTGCGAC	AGCCCTCGCC	TCCGACCCCA	1000
CCCGTTGCGC TGGAAGGAAT TCGTGCGGCG CCTAGGGCTG AGCGACCACG 1150 AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCGCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCG ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCGGCC CCGCCGCCT CCCGCCCGCG 1350	TCCCCAACCC	CCTTCAGAAG	TGGGAGGACA	GCGCCCACAA	GCCACAGAGC	1050
AGATCGATCG GCTGGAGCTG CAGAACGGGC GCTGCCTGCG CGAGGCGCAA 1200 TACAGCATGC TGGCGACCTG GAGGCGGCC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCGGCC CCGCCGCCT CCCGCCCGCG 1350	CTAGACACTG	ATGACCCCGC	GACGCTGTAC	GCCGTGGTGG	AGAACGTGCC	1100
TACAGCATGC TGGCGACCTG GAGGCGGCGC ACGCCGCGGC GCGAGGCCAC 1250 GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCGGCC CCGCCGCCT CCCGCCCGCG 1350	CCCGTTGCGC	TGGAAGGAAT	TCGTGCGGCG	CCTAGGGCTG	AGCGACCACG	1150
GCTGGAGCTG CTGGGACGCG TGCTCCGCGA CATGGACCTG CTGGGCTGCC 1300 TGGAGGACAT CGAGGAGGCG CTTTGCGGCC CCGCCGCCT CCCGCCCGCG 1350	AGATCGATCG	GCTGGAGCTG	CAGAACGGGC	GCTGCCTGCG	CGAGGCGCAA	1200
TGGAGGACAT CGAGGAGGCG CTTTGCGGCC CCGCCGCCT CCCGCCCGCG 1350	TACAGCATGC	TGGCGACCTG	GAGGCGGCGC	ACGCCGCGGC	GCGAGGCCAC	1250
	GCTGGAGCTG	CTGGGACGCG	TGCTCCGCGA	CATGGACCTG	CTGGGCTGCC	1300
ርርር <u>ን</u> ርጥርጥር ጥር <u>ን</u> ርንጥርን	TGGAGGACAT	CGAGGAGGCG	CTTTGCGGCC	CCGCCGCCCT	CCCGCCCGCG	1350
CCCAGTOTIC TCAGATGA	CCCAGTCTTC	TCAGATGA				1365

(3) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 483 bases
- (B) TYPE:nucleic acid
- (C) STRANDEDNESS:single
- (D) TOPOLOGY:linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

GATAGTGTGT	GTCCCCAAGG	AAAATATATC	CACCCTCAAA	ATAATTCGAT	50
TTGCTGTACC	AAGTGCCACA	AAGGAACCTA	CTTGTACAAT	GACTGTCCAG	100
GCCCGGGGCA	GGATACGGAC	TGCAGGGAGT	GTGAGAGCGG	CTCCTTCACC	150
GCTTCAGAAA	ACCACCTCAG	ACACTGCCTC	AGCTGCTCCA	AATGCCGAAA	200
GGAAATGGGT	CAGGTGGAGA	TCTCTTCTTG	CACAGTGGAC	CGGGACACCG	250
TGTGTGGCTG	CAGGAAGAAC	CAGTACCGGC	ATTATTGGAG	TGAAAACCTT	300

TTCCAGTGCT	TCAATTGCAG	CCTCTGCCTC	AATGGGACCG	TGCACCTCTC	350
CTGCCAGGAG	AAACAGAACA	CCGTGTGCAC	CTGCCATGCA	GGTTTCTTTC	400
TAAGAGAAAA	CGAGTGTGTC	TCCTGTAGTA	ACTGTAAGAA	AAGCCTGGAG	450
TGCACGAAGT	TGTGCCTACC	CCAGATTGAG	AAT		483

(4) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:455 amino acids
- (B) TYPE:polypeptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Gly	Leu	Ser	Thr 5	Val	Pro	Asp	Leu	Leu 10	Leu	Pro	Leu	Val	Leu 15
Leu Glu	Leu	Leu	Val 20	Gly	Ile	Tyr	Pro	Ser 25	Gly	Val	Ile	Gly	Leu 30
Val Pro	His	Leu	Gly 35	Asp	Arg	Glu	Lys	Arg 40	Asp	Ser	Val	Cys	Pro 45
Gln Gly	Lys	Tyr	Ile 50	His	Pro	Gln	Asn	Asn 55	Ser	Ile	Cys	Cys	Thr 60
Lys Cys			65		_		_	70	_	_		_	75
Gly Gln	_		80	_				85		_			90
Ala Ser			95				-	100		-		-	105
Arg Lys			110					115		_			Asp 120
Arg Asp			125	_	_		_	130		_			Tyr 135
Trp Ser			140			_		145	-			-	150
Asn Gly	Thr	Val	His 155	Leu	Ser	Cys	Gln	Glu 160	Lys	Gln	Asn	Thr	Val 165
Cys Thr	_		170	_				175				_	Val 180
Ser Cys	Ser	Asn	Cys 185	Lys	Lys	Ser	Leu	Glu 190	Cys	Thr	Lys	Leu	Cys 195
Leu Pro	Gln	Ile	Glu 200	Asn	Val	Lys	Gly	Thr 205	Glu	Asp	Ser	Gly	Thr 210
Thr Val	Leu	Leu	Pro 215	Leu	Val	Ile	Phe	Phe 220	Gly	Leu	Cys	Leu	Leu 225
Ser Leu	Leu	Phe	Ile 230	Gly	Leu	Met	Tyr	Arg 235	Tyr	Gln	Arg	Trp	Lys 240
Ser Lys	Leu	Tyr	Ser 245	Ile	Val	Cys	Gly	Lys 250	Ser	Thr	Pro	Glu	Lys 255

Glu Gly Glu Leu Glu Gly Thr Thr Thr Lys Pro Leu Ala Pro Asn Pro Ser Phe Ser Pro Thr Pro Gly Phe Thr Pro Thr Leu Gly Phe Ser Pro Val Pro Ser Ser Thr Phe Thr Ser Ser Ser Thr Tyr Thr Pro Gly Asp Cys Pro Asn Phe Ala Ala Pro Arg Arg Glu Val Ala Pro Pro Tyr Gln Gly Ala Asp Pro Ile Leu Ala Thr Ala Leu Ala Ser Asp Pro Ile Pro Asn Pro Leu Gln Lys Trp Glu Asp Ser Ala His Lys Pro Gln Ser Leu Asp Thr Asp Asp Pro Ala Thr Leu Tyr Ala Val Val Glu Asn Val Pro Pro Leu Arg Trp Lys Glu Phe Val Arg Arg Leu Gly Leu Ser Asp His Glu Ile Asp Arg Leu Glu Leu Gln Asn Gly Arg Cys Leu Arg Glu Ala Gln Tyr Ser Met Leu Ala Thr Trp Arg Arg Thr Pro Arg Arg Glu Ala Thr Leu Glu Leu Leu Gly Arg Val Leu Arg Asp Met Asp Leu Leu Gly Cys Leu Glu Asp Ile Glu Glu Ala Leu Cys Gly Pro Ala Ala Leu Pro Pro Ala Pro Ser Leu Leu Arq

- (5) INFORMATION FOR SEQ ID NO:4:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 161 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
- Asp Ser Val Cys Pro Gln Gly Lys Tyr Ile His Pro Gln Asn Asn
 5 10 15

 Ser Ile Cys Cys Thr Lys Cys His Lys Gly Thr Tyr Leu Tyr Asn
 20 25 30

 Asp Cys Pro Gly Pro Gly Gln Asp Thr Asp Cys Arg Glu Cys Glu
 35 40 45

 Ser Gly Ser Phe Thr Ala Ser Glu Asn His Leu Arg His Cys Leu
 50 55 60

 Ser Cys Ser Lys Cys Arg Lys Glu Met Gly Gln Val Glu Ile Ser

Ser	Cys	Thr	Val	Asp	Arg	Asp	Thr	Val	Cys	Gly	Cys	Arg	Lys	Asn
				80					85					90
Gln	Tyr	Arg	His	Tyr	Trp	Ser	Glu	Asn	Leu	Phe	Gln	Cys	Phe	Asn
				95					100					105
Cys	Ser	Leu	Cys	Leu	Asn	Gly	Thr	Val	His	Leu	Ser	Cys	Gln	Glu
				110					115					120
Lys	Gln	Asn	Thr	Val	Cys	Thr	Cys	His	Ala	Gly	Phe	Phe	Leu	Arg
				125					130					135
Glu	Asn	Glu	Cys	Val	Ser	Cys	Ser	Asn	Cys	Lys	Lys	Ser	Leu	Glu
				140					145					150
Cys	Thr	Lys	Leu	Cys	Leu	Pro	Gln	Ile	Glu	Asn				
				155					160					

- (6) INFORMATION FOR SEQ ID NO:5:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:157 bases
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS:single
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

CAGGGGAAA	ATATTCACCC	TCAAATAAT	TCGATTTGCT	GTACCAAGTG	50
CCACAAAGG	AAACTACTTG	TACAATGAC	TGTCCAGGCC	CGGGGCAGGA	100
TACGGACTG	CAGGGAGTGT	GAGAGCGGC	TCCTTCACAG	CCTCAGAAAA	150
CAACAAG					157

- (7) INFORMATION FOR SEQ ID NO:6:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:13 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Asp Ser Val Xaa Pro Gln Gly Lys Tyr Ile His Pro Gln 5 10

- (8) INFORMATION FOR SEQ ID NO:7:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:11 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Glu Asn Glu Cys Val Ser Cys Ser Asn Cys Lys
5 10

- (9) INFORMATION FOR SEQ ID NO:8:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:12 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
- Glu Asn Glu Cys Val Ser Cys Ser Asn Cys Lys Lys 5
- (10) INFORMATION FOR SEQ ID NO:9:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 13 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
- Tyr Ile His Pro Gln Xaa Asn Ser Ile Xaa Xaa Xaa Lys
 5 10
- 11) INFORMATION FOR SEQ ID NO:10:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 14 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Glu Cys Glu Ser Gly Ser Phe Thr Ala Ser Glu Asn Asn Lys
 5 10
- (12) INFORMATION FOR SEQ ID NO:11:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 15 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
- Lys Glu Met Gly Gln Val Glu Ile Ser Ser Cys Thr Val Asp Arg
 5 10 15

- (13) INFORMATION FOR SEQ ID NO:12:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:13 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:
- Gly Thr Tyr Ley Tyr Asn Asp Cys Pro Gly Pro Gly Gln
 5 10
- (14) INFORMATION FOR SEQ ID NO:13:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 13 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
- Glu Met Gly Gln Val Glu Ile Ser Xaa Xaa Xaa Val Asp
 5 10
- (15) INFORMATION FOR SEQ ID NO:14:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 20 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Lys Glu Met Gly Gln Val Glu Ile Ser Ser Cys Thr Val Asp Arg
5 10 15
Asp Thr Val Cys Gly
20

- (16) INFORMATION FOR SEQ ID NO:15:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 19 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Tyr Ile His Pro Gln Xaa Asn Ser Ile Cys Cys Thr Lys Cys His $5 \hspace{1.5cm} 10 \hspace{1.5cm} 15$ Lys Gly Xaa Tyr

- (17) INFORMATION FOR SEQ ID NO:16:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 18 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Gly Thr Tyr Leu Tyr Asn Asp Cys Pro Gly Pro Gly Gln Asp Thr
5 10 15
Xaa Xaa Arg

- (18) INFORMATION FOR SEQ ID NO:17:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 14 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:
- Gln Asn Thr Val Cys Thr Xaa His Ala Gly Phe Phe Leu Arg
 5 10
- (19) INFORMATION FOR SEQ ID NO:18:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 14 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Ser Leu Glu Cys Thr Lys Leu Cys Leu Pro Gln Ile Glu Asn 5 10

- (20) INFORMATION FOR SEQ ID NO:19:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 13 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Asp Ser Val Cys Pro Gln Gly Lys Tyr Ile His Pro Gln
5 10

(21) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
CAAGGTAAAT ATATTCATCC	20
(22) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CAGGGTAAGT ACATCCATCC	20
(23) INFORMATION FOR SEQ ID NO:22:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
CAAGGTAAAT ATATACATCC	20
(24) INFORMATION FOR SEQ ID NO:23:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
CAAGGCAAAT ATATTCATCC	20

(25) INFORMATION FOR SEQ ID NO:24:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
CAGGGCAAGT ACATCCACCC	20
(26) INFORMATION FOR SEQ ID NO:25:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
CAAGGCAAAT ATATACATCC	20
(27) INFORMATION FOR SEQ ID NO:26:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
CAAGGAAAAT ATATTCATCC	20
(28) INFORMATION FOR SEQ ID NO:27:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SECTENCE DESCRIPTION: SEC ID NO:27:	

CAGGGAAAGT ACATCCACCC	20
(29) INFORMATION FOR SEQ ID NO:28:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
CAAGGAAAAT ATATACATCC	20
(30) INFORMATION FOR SEQ ID NO:29:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
CAAGGGAAAT ATATTCATCC	20
(31) INFORMATION FOR SEQ ID NO:30:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
CAGGGGAAGT ACATCCACCC	20
(31) INFORMATION FOR SEQ ID NO:31:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:20 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	

CAAGGGAAAT ATATACATCC	20
(33) INFORMATION FOR SEQ ID NO:32:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:14 amino acids(B) TYPE:polypeptide	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
Glu Cys Gly Ser Gly Ser Phe Thr Ala Ser Glu Asn Asn Lys 5 10	
(34) INFORMATION FOR SEQ ID NO:33:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:14 amino acids(B) TYPE:polypeptide	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
Glu Cys Gly Ser Gly Ser Phe Thr Ala Ser Cys Asn Asn Lys 5 10	
(35) INFORMATION FOR SEQ ID NO:34:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
AAATGACGGA GACTCTTGTT GTTCCTAGGG	30
(36) INFORMATION FOR SEQ ID NO:35:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

AAGTGGCGTA GTCTTTGTT GTTCCTAGGG	30
(37) INFORMATION FOR SEQ ID NO:36:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 30 bases	
(B) TYPE:nucleic acid	
(C) STRANDEDNESS:single	
(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
AAATGTCGGA GACTCTTGTT GTTCCTAGGG	30
(38) INFORMATION FOR SEQ ID NO:37:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 30 bases	
(B) TYPE:nucleic acid	
(C) STRANDEDNESS:single	
(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:	
AAATGACGGT CACTCTTGTT GTTCCTAGGG	30
(39) INFORMATION FOR SEQ ID NO:38:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 30 bases	
(B) TYPE:nucleic acid	
(C) STRANDEDNESS:single	
(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:	
AAGTGGCGTT CTCTTTGTT GTTCCTAGGG	30
(40) INFORMATION FOR SEQ ID NO:39:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 30 bases	
(B) TYPE:nucleic acid	
(C) STRANDEDNESS:single	
(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:	

AAATGTCGGT CACTCTTGTT GTTCCTAGGG	30
(41) INFORMATION FOR SEQ ID NO:40:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:	
AAATGACGGA GAACATTGTT GTTCCTAGGG	30
(42) INFORMATION FOR SEQ ID NO:41:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:	
AAGTGGCGTA GTACTTTGTT GTTCCTAGGG	30
(43) INFORMATION FOR SEQ ID NO:42:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:	
AAATGTCGGA GAACATTGTT GTTCCTAGGG	30
(44) INFORMATION FOR SEQ ID NO:43:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
AAATGACGGT CAACATTGTT GTTCCTAGGG	0
(45) INFORMATION FOR SEQ ID NO:44:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
AAGTGGCGTT CTACTTTGTT GTTCCTAGGG	0
(46) INFORMATION FOR SEQ ID NO:45:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:	
AAATGTCGGT CAACATTGTT GTTCCTAGGG 3	0
(47) INFORMATION FOR SEQ ID NO:46:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:51 amino acids(B) TYPE:polypeptide	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:	
Gln Gly Lys Tyr Ile His Pro Gln Asn Asn Ser Ile Cys Cys Th	ır .5
Lys Cys His Lys Gly Thr Tyr Ley Tyr Asn Asp Cys Pro Gly Pr	
Gly Gln Asp Thr Asp Cys Arg Gly Cys Glu Ser Gly Ser Phe Th	r :5
Ala Ser Glu Asn Asn Lys 50	
(48) INFORMATION FOR SEQ ID NO:47:	

(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:158 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
CAGGGGAAAT ATATTCACCC TCAAAATAAT TCGATTTCGT GTACCAAGTC GCACAAAGGA ACCTACTTGT ACAATGACTG TCCAGGCCCG GGGCAGGATA CGGACTGCAG GGAGTGTGAG AGCGGCTCCT TCACAGCCTC AGAAAACAAC AAGGATCC	100
(49) INFORMATION FOR SEQ ID NO:48:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:26 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
GGAATTCAGC CTGAATGGCG AATGGG	26
(50) INFORMATION FOR SEQ ID NO:49:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:25 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:	
CCTCGAGCGT TGCTGGCGTT TTTCC	25
(51) INFORMATION FOR SEQ ID NO:50:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:23 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:	

GGTCGACATT GATTATTGAC TAG	23
(52) INFORMATION FOR SEQ ID NO:51:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:23 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:	
GGAATTCCCT AGGAATACAG CGG	23
(53) INFORMATION FOR SEQ ID NO:52:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:18 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:	
GGCAAGGGCA GCAGCCGG	18
(54) INFORMATION FOR SEQ ID NO:53:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:53 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:	
AGCTTCTGCA GGTCGACATC GATGGATCGG TACCTCGAGC GGCCGCGAAT	50 53
(55) INFORMATION FOR SEQ ID NO:54:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH:54 bases (B) TYPE:nucleic acid (C) STRANDEDNESS:single (D) TOPOLOGY:linear 	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:	
CTAGAGAATT CGCGGCCGCT CGAGGTACCG GATCCATCGA TGTCGACCTG CAGA	50 54
(56) INFORMATION FOR SEQ ID NO:55:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:63 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:	
AGCTCTAGAG ATTCGCGGCC GCTCGAGGTA CCGGATCCAT CGATGTCGAC CTGCAGAAGC TTG	50 63
(57) INFORMATION FOR SEQ ID NO:56:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:64 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:	
CTAGCAAGCT TCTGCAGGTC GACATCGATG GATCCGGTAC CTCGAGCGGC CGCGAATTCT CTAG	50 64
(58) INFORMATION FOR SEQ ID NO:57:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:25 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:	
CAGGATCCGA GTCTCAACCC TCAAC	25
(59) INFORMATION FOR SEQ ID NO:58:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 43 bases	

(C) STRANDEDNESS:single	
(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:	
GGGAATTCCT TATCAATTCT CAATCTGGGG TAGGCACAAC TTC	43
(60) INFORMATION FOR SEQ ID NO:59:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:81 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:	
CACAGTCGAC TTACATTTGC TTCTGACACA ACTGTGTTCA CTAGCAACCT CAAACAGACA CCATGGGCCT CTCCACCGTG C	50 81
(61) INFORMATION FOR SEQ ID NO:60:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:17 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:	
GAGGCTGCAA TTGAAGC	17
(62) INFORMATION FOR SEQ ID NO:61:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:17 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:	
ATTCGTGCGG CGCCTAG	17
(63) INFORMATION FOR SEO ID NO:62:	

(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:17 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:		
GTCGGTAGCA CCAAGGA	17	
(64) INFORMATION FOR SEQ ID NO:63:		
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:17 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:		
GTTTTCCCAG TCACGAC	17	
(65) INFORMATION FOR SEQ ID NO:64:		
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:30 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:		
GTCCAATTAT GTCACACC	18	
(66) INFORMATION FOR SEQ ID NO:65:		
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH:1334 bases(B) TYPE:nucleic acid(C) STRANDEDNESS:single(D) TOPOLOGY:linear		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:		
GAATTCTCTG GACTGAGGCT CCAGTTCTGG CCTTTGGGGT TCAAGATCAC TGGGACCAGG CCGTGATCTC TATGCCCGAG TCTCAACCCT CAACTGTCAC		50 100

	ragg(CGAG			200
TAG	CTGT	CTG (ЭC		•					212
							CTG Leu			257
							GGG Gly			302
							GAT Asp			347
							TCG Ser			392
							GAC Asp			437
							AGC Ser			482
							AGC Ser			527
			-				TCT Ser			572
							CAG Gln			617
							TGC Cys			662
							AAA Lys			707

				155					160					165	
										GAA Glu					752
										TGC Cys					797
										GAG Glu					842
										GGT Gly					887
										TAC Tyr					932
										TCG Ser					977
										CCC Pro					1022
					Thr		Gly	Phe		CCC Pro					1067
										AGC Ser					1112
										CGC Arg					1157
										GCG Ala					1202
TCC	GAC	CCC	ATC	CCC	AAC	CCC	CTT	CAG	AAG	TGG	GAG	GAC	AGC	GCC	1247

(67) INFORMATION FOR SEQ ID NO:66:

365

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 371 amino acids
- (B) TYPE:polypeptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:

Met Gly Leu Ser Thr Val Pro Asp Leu Leu Pro Leu Val Leu Leu Glu Leu Leu Val Gly Ile Tyr Pro Ser Gly Val Ile Gly Leu Val Pro His Leu Gly Asp Arg Glu Lys Arg Asp Ser Val Cys Pro 35 40 Gln Gly Lys Tyr Ile His Pro Gln Asn Asn Ser Ile Cys Cys Thr 50 Lys Cys His Lys Gly Thr Tyr Leu Tyr Asn Asp Cys Pro Gly Pro 75 Gly Gln Asp The Asp Cys Arg Glu Cys Gly Ser Gly Ser Phe Thr Ala Ser Glu Asn His Leu Arq His Cys Leu Ser Cys Ser Lys Cys 105 Arg Lys Glu Met Gly Gln Val Glu Ile Ser Ser Cys Thr Val Asp 110 115 120 Arg Asp Thr Val Cys Gly Cys Arg Lys Asn Gln Tyr Arg His Tyr 125 130 135 Trp Ser Glu Asn Leu Phe Gln Cys Phe Asn Cys Ser Leu Cys Leu 140 145 150 Asn Gly Thr Val His Leu Ser Cys Gln Glu Lys Gln Asn Thr Val 155 160 Cys Thr Cys His Ala Gly Phe Phe Leu Arg Glu Asn Glu Cys Val 175 180 Ser Cys Ser Asn Cys Lys Lys Ser Leu Glu Cys Thr Lys Leu Cys 185 190 Leu Pro Gln Ile Glu Asn Val Lys Gly Thr Glu Asp Ser Gly Thr

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200
                                     205
                                                          210
Thr Val Leu Leu Pro Leu Val Ile Phe Phe Gly Leu Cys Leu Leu
                215
                                     220
                                                          225
Ser Leu Leu Phe Ile Gly Leu Met Tyr Arg Tyr Gln Arg Trp Lys
                                     235
                                                          240
Ser Lys Leu Tyr Ser Ile Val Cys Gly Lys Ser Thr Pro Glu Lys
                245
                                     250
                                                          255
Glu Gly Glu Leu Glu Gly Thr Thr Thr Lys Pro Leu Ala Pro Asn
                260
                                     265
Pro Ser Phe Ser Pro Thr Pro Gly Phe Thr Pro Thr Leu Gly Phe
                275
                                     280
Ser Pro Val Pro Ser Ser Thr Phe Thr Ser Ser Ser Thr Tyr Thr
                290
                                     295
Pro Gly Asp Cys Pro Asn Phe Ala Ala Pro Arg Arg Glu Val Ala
                                     310
                                                          315
Pro Pro Tyr Gln Gly Ala Asp Pro Ile Leu Ala Thr Ala Leu Ala
                320
                                     325
                                                          330
Ser Asp Pro Ile Pro Asn Pro Leu Gln Lys Trp Glu Asp Ser Ala
                335
                                     340
                                                          345
His Lys Pro Gln Ser Leu Asp Thr Asp Asp Pro Ala Thr Leu Tyr
                350
                                     355
Ala Val Val Glu Asn Val Pro Pro Leu Arg Trp
                365
                                     370
```

(68) INFORMATION FOR SEQ ID NO:67:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 6464 bases
- (B) TYPE:nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:

TCGACATTGA	TTATTGACTA	GTTATTAATA	GTAATCAATT	ACGGGGTCAT	50
TAGTTCATAG	CCCATATATG	GAGTTCCGCG	TTACATAACT	TACGGTAAAT	100
GGCCCGCCTC	GCTGACCGCC	CAACGACCCC	CGCCCATTGA	CGTCAATAAT	150
GACGTATGTT	CCCATAGTAA	CGCCAATAGG	GACTTTCCAT	TGACGTCAAT	200
GGGTGGAGTA	TTTACGGTAA	ACTGCCCACT	TGGCAGTACA	TCAAGTGTAT	250
CATATGCCAA	GTACGCCCCC	TATTGACGTC	AATGACGGTA	AATGGCCCGC	300
CTGGCATTAT	GCCCAGTACA	TGACCTTATG	GGACTTTCCT	ACTIGGCAGT	350
ACATCTACGT	ATTAGTCATC	GCTATTACCA	TGGTGATGCG	GTTTTGGCAG	400
TACATCAATG	GGCGTGGATA	GCGGTTTGAC	TCACGGGGAT	TTCCAAGTCT	450
CCACCCCATT	GACGTCAATG	GGAGTTTGTT	TTGGCACCAA	AATCAACGGG	500
ACTTTCCAAA	ATGTCGTAAC	AACTCCGCCC	CATTGACGCA	AATGGGCGGT	550
AGGCGTGTAC	GGTGGGAGGT	CTATATAAGC	AGAGCTCTCT	GGCTAACTAG	600

AGAACCCACT	GCTTAACTGG	CTTATCGAAA	TTAATACGAC	TCACTATAGG	650
GAGACCCAAG	CTTCTGCAGG	TCGACATCGA	TGGATCCGGT	ACCTCGAGCG	700
CGAATTCTCT	AGAGGATCTT	TGTGAAGGAA	CCTTACTTCT	GTGGTGTGAC	750
ATAATTGGAC	AAACTACCTA	CAGAGATTTA	AAGCTCTAAG	GTAAATATAA	800
AATTTTTAAG	TGTATAATGT	GTTAAACTAC	TGATTCTAAT	TGTTTGTGTA	850
TTTTAGATTC	CAACCTATGG	AACTGATGAA	TGGGAGCAGT	GGTGGAATGC	900
CTTTAATGAG	GAAAACCTGT	TTTGCTCAGA	AGAAATGCCA	TCTAGTGATG	950
ATGAGGCTAC	TGCTGACTCT	CAACATTCTA	CTCCTCCAAA	AAAGAAGAGA	1000
AAGGTAGAAG	ACCCCAAGGA	CTTTCCTTCA	GAATTGCTAA	GTTTTTTGAG	1050
TCATGCTGTG	TTTAGTAATA	GAACTCTTGC	TTGCTTTGCT	ATTTACACCA	1100
CAAAGGAAAA	AGCTGCACTG	CTATACAAGA	AAATTATGGA	AAAATATTTG	1150
ATGTATAGTG	CCTTGACTAG	AGATCATAAT	CAGCCATACC	ACATTTGTAG	1200
AGGTTTTACT	TGCTTTAAAA	AACCTCCCAC	ACCTCCCCCT	GAACCTGAAA	1250
CATAAAATGA	ATGCAATTGT	TGTTGTTAAC	TTGTTTATTG	CAGCTTATAA	1300
TGGTTACAAA	TAAAGCAATA	GCATCACAAA	TTTCACAAAT	AAAGCATTTT	1350
TTTCACTGCA	TTCTAGTTGT	GGTTTGTCCA	AACTCATCAA	TGTATCTTAT	1400
CATGTCTGGA	TCAATTCTGA	GAAACTAGCC	TTAAAGACAG	ACAGCTTTGT	1450
TCTAGTCAGC	CAGGCAAGCA	TATGTAAATA	AAGTTCCTCA	GGGAACTGAG	1500
GTTAAAAGAT	GTATCCTGGA	CCTGCCAGAC	CTGGCCATTC	ACGTAAACAG	1550
AAGATTCCGC	CTCAAGTTCC	GGTTAACAAC	AGGAGGCAAC	GAGATCTCAA	1600
ATCTATTACT	TCTAATCGGG	TAATTAAAAC	CTTTCAACTA	AAACACGGAC	1650
CCACGGATGT	CACCCACTTT	TCCTTCCCCG	GCTCCGCCCT	TCTCAGTACT	1700
CCCCACCATT	AGGCTCGCTA	CTCCACCTCC	ACTTCCGGGC	GCGACACCCA	1750
CGTGCCCTCT	CCCACCCGAC	GCTAACCCCG	CCCCTGCCCG	TCTGACCCCG	1800
CCCACCACCT	GGCCCCGCCC	CGTTGAGGAC	AGAAGAAACC	CCGGGCAGCC	1850
GCAGCCAAGG	CGGACGGGTA	GACGCTGGGG	GCGCTGAGGA	GTCGTCCTCT	1900
ACCTTCTCTG	CTGGCTCGGT	GGGGGACGCG	GTGGATCTCA	GGCTTCCGGA	1950
AGACTGGAAG	AACCGGCTCA	GAACCGCTTG	TCTCCGCGGG	GCTTGGGCGG	2000
CGGAAGAATG	GCCGCTAGAC	GCGGACTTGG	TGCGAGGCAT	CGCAGGATGC	2050
AGAAGAGCAA	GCCCGCCGGG	AGCGCGCGC	TGTACTACCC	CGCGCCTGGA	2100
GCGGCCACGC	CGGACTGGGC	GGGGCCGGCC	TGGTGGAGGC	GGAGTCTGAC	2150
CTCGTGGAGG	CGGGGCCTCT	GATGTTCAAA	TAGGATGCTA	GGCTTGTTGA	2200
GGCGTGGCCT	CCGATTCACA	AGTGGGAAGC	AGCGCCGGGC	GACTGCAATT	2250
TCGCGCCAAA	CTTGGGGGAA	GCACAGCGTA	CAGGCTGCCT	AGGTGATCGC	2300
TGCTGCTGTC	ATGGTTCGAC	CGCTGAACTG	CATCGTCGCC	GTGTCCCAGA	2350
ATATGGGCAT	CGGCAAGAAC	GGAGACCTTC	CCTGGCCAAT	GCTCAGGTAC	2400
TGGCTGGATT	GGGTTAGGGA	AACCGAGGCG	GTTCGCTGAA	TCGGGTCGAG	2450
CACTTGGCGG	AGACGCGCGG	GCCAACTACT	TAGGGACAGT	CATGAGGGGT	2500
AGGCCCGCCG	GCTGCTGCCC	TTGCCCATGC	CCGCGGTGAT	CCCCATGCTG	2550
TGCCAGCCTT	TGCCCAGAGG	CGCTCTAGCT	GGGAGCAAAG	TCCGGTCACT	2600
GGGCAGCACC	ACCCCCGGA	${\tt CTTGCATGGG}$	${\tt TAGCCGCTGA}$	GATGGAGCCT	2650
GAGCACACGT	GACAGGGTCC	CTGTTAACGC	AGTGTTTCTC	TAACTTTCAG	2700
GAACGAGTTC	AAGTACTTCC	AAAGAATGAC	CACCACCTCC	TCAGTGGAAG	2750
GTAAACAGAA	CCTGGTGATT	${\tt ATGGGCCGGA}$	AAACCTGGTT	$\mathtt{CTCCATTCCT}$	2800
GAGAAGAATC	GACCTTTAAA	GGACAGAATT	AATATAGTTC	TCAGTAGAGA	2850
GCTCAAGGAA	CCACCACAAG	GAGCTCATTT	TCTTGCCAAA	AGTCTGGACC	2900

3 maaamma 3 3	3 CEEE EE C 3 3	G3 3 GG3 G3 GE			
				AGTGGACATG	
		TTCCGTTTAC			3000
	AGACTCTTTG	TGACAAGGAT	CATGCAGGAA	TTTGAAAGTG	3050
ACACGTTCTT	CCCAGAAATT	GATTTGGAGA		TCTCCCAGAG	3100
TACCCAGGGG			GAAAAAGGCA	TCAAGTATAA	3150
ATTTGAAGTC		AAGGCTAACA		TGCTGATTGA	3200
CTTCAAGTTC		CTCCTAAAAT	TATGCATTTT	TACAAGACCA	3250
	GTTGGCTTTA				3300
	TCCCCAAAGT			ATTCTAAACA	3350
ATTAGAATTA	TTTTCATTTT	CATTAGTCTA	ACCAGGTTAT	ATTAAATATA	3400
CTTTAAGAAA	CACCATTTGC	CATAAAGTTC	TCAATGCCCC	TCCCATGCAG	3450
CCTCAAGTGG	CTCCCCAGCA	GATGCATAGG	GTAGTGTGTG	TACAAGAGAC	3500
CCCAAAGACA	TAGAGCCCCT	GAGAGCATGA	GCTGATATGG	GGGCTCATAG	3550
AGATAGGAGC	TAGATGAATA	AGTACAAAGG	GCAGAAATGG	GTTTTAACCA	3600
GCAGAGCTAG	AACTCAGACT	TTAAAGAAAA	TTAGATCAAA	GTAGAGACTG	3650
AATTATTCTG	CACATCAGAC	TCTGAGCAGA	GTTCTGTTCA	CTCAGACAGA	3700
AAATGGGTAA	ATTGAGAGCT	GGCTCCATTG	TGCTCCTTAG	AGATGGGAGC	3750
AGGTGGAGGA	TTATATAAGG	TCTGGAACAT	TTAACTTCTC	CGTTTCTCAT	3800
CTTCAGTGAG	ATTCCAAGGG	ATACTACAAT	TCTGTGGAAT	GTGTGTCAGT	3850
TAGGGTGTGG	AAAGTCCCCA	GGCTCCCCAG	CAGGCAGAAG	TATGCAAAGC	3900
ATGCATCTCA	ATTAGTCAGC	AACCAGGTGT	GGAAAGTCCC	CAGGCTCCCC	4000
AGCAGGCAGA	AGTATGCAAA	GCATGCATCT	CAATTAGTCA	GCAACCATAG	4050
TCCCGCCCCT	AACTCCGCCC	ATCCCGCCCC	TAACTCCGCC	CAGTTCCGCC	4100
CATTCTCCGC	CCCATGGCTG	ACTAATTTTT	TTTATTTATG	CAGAGGCCGA	4150
GGCGCCTCTG	AGCTATTCCA	GAAGTAGTGA	GGAGGCTTTT	TTGGAGGCCT	4200
AGGCTTITGC	AAAAAAGCTA	ATTCAGCCTG	AATGGCGAAT	GGGACGCGCC	4250
CTGTAGCGGC	GCATTAAGCG	CGGCGGGTGT	GGTGGTTACG	CGCAGCGTGA	4255
CCGCTACACT	TGCCAGCGCC	CTAGCGCCCG	CTCCTTTCGC	TTTCTTCCCT	4300
TCCTTTCTCG	CCACGTTCGC	CGGCTTTCCC	CGTCAAGCTC	TAAATCGGGG	4350
GCTCCCTTTA	GGGTTCCGAT	TTAGTGCTTT	ACGGCACCTC	GACCCCAAAA	4400
ACTTGATTAG	GGTGATGGTT	CACGTAGTGG	GCCATCGCCC	TGATAGACGG	4450
TTTTTCGCCC	TTTGACGTTG	GAGTCCACGT	TCTTTAATAG	TGGACTCTTG	4500
TTCCAAACTG	GAACAACACT	CAACCCTATC	TCGGTCTATT	CTTTTGATTT	4550
				GAGCTGATTT	4600
AACAAAAATT	TAACGCGAAT	TTTAACAAAA	TATTAACGTT	TACAATTTCA	4650
				GTTTATTTTT	
CTAAATACAT	TCAAATATGT	ATCCGCTCAT	GAGACAATAA	CCCTGATAAA	4750
				ACATTTCCGT	
				TTTTTGCTCA	
				TTGGGTGCAC	
				CCTTGAGAGT	
				AAGTTCTGCT	
				CAACTCGGTC	
				ACCAGTCACA	
				GCAGTGCTGC	
				ACAACGATCG	
				GGATCATGTA	

ACTCGCCTTG	ATCGTTGGGA	ACCGGAGCTG	AATGAAGCCA	TACCAAACGA	5300
CGAGCGTGAC	ACCACGATGC	CTGTAGCAAT	GGCAACAACG	TTGCGCAAAC	5350
TATTAACTGG	CGAACTACTT	ACTCTAGCTT	CCCGGCAACA	ATTAATAGAC	5400
TGGATGGAGG	CGGATAAAGT	TGCAGGACCA	CTTCTGCGCT	CGGCCCTTCC	5450
GGCTGGCTGG	TTTATTGCTG	ATAAATCTGG	AGCCGGTGAG	CGTGGGTCTC	5500
GCGGTATCAT	TGCAGCACTG	GGGCCAGATG	GTAAGCCCTC	CCGTATCGTA	5550
GTTATCTACA	CGACGGGGAG	TCAGGCAACT	ATGGATGAAC	GAAATAGACA	5600
GATCGCTGAG	ATAGGTGCCT	CACTGATTAA	GCATTGGTAA	CTGTCAGACC	5650
AAGTTTACTC	ATATATACTT	TAGATTGATT	TAAAACTTCA	TTTTTAATTT	5700
AAAAGGATCT	AGGTGAAGAT	CCTTTTTGAT	AATCTCATGA	CCAAAATCCC	5750
TTAACGTGAG	TTTTCGTTCC	ACTGAGCGTC	AGACCCCGTA	GAAAAGATCA	5800
AAGGATCTTC	TTGAGATCCT	TTTTTTCTGC	GCGTAATCTG	CTGCTTGCAA	5850
ACAAAAAAAC	CACCGCTACC	AGCGGTGGTT	TGTTTGCCGG	ATCAAGAGCT	5900
ACCAACTCTT	TTTCCGAAGG	TAACTGGCTT	CAGCAGAGCG	CAGATACCAA	6000
ATACTGTCCT	TCTAGTGTAG	CCGTAGTTAG	GCCACCACTT	CAAGAACTCT	6050
GTAGCACCGC	CTACATACCT	CGCTCTGCTA	ATCCTGTTAC	CAGTGGCTGC	6100
TGCCAGTGGC	GATAAGTCGT	GTCTTACCGG	GTTGGACTCA	AGACGATAGT	6150
TACCGGATAA	GGCGCAGCGG	TCGGGCTGAA	CGGGGGGTTC	GTGCACACAG	6200
CCCAGCTTGG	AGCGAACGAC	CTACACCGAA	CTGAGATACC	TACAGCGTGA	6250
GCATTGAGAA	AGCGCCACGC	TTCCCGAAGG	GAGAAAGGCG	GACAGGTATC	6300
CGGTLAGCGG	CAGGGTCGGA	ACAGGAGAGC	GCACGAGGGA	GCTTCCAGGG	6350
GGAAACGCCT	GGTATCTTTA	TAGTCCTGTC	GGGTTTCGCC	ACCTCTGACT	6400
TGAGCGTCGA	TTTTTGTGAT	GCTCGTCAGG	GGGGCGGAGC	CTATGGAAAA	6450
ACGCCAGCAA	CGCC				6464

(69) INFORMATION FOR SEQ ID NO:68:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:2173 bases
- (B) TYPE:nucleic acid
- (C) STRANDEDNESS:single
- (D) TOPOLOGY:linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:

GAATTCCTTT TCTCCGAGTT TTCTGAACTC TGGCTCATGA TCGGGCTTAC	50
TGGATACGAG AATCCTGGAG GACCGTACCC TGATTTCCAT CTACCTCTGA	100
CTTTGAGCCT TTCTAACCCG GGGCTCACGC TGCCAACACC CGGGCCACCT	150
GGTCCGATCG TCTTACTTCA TTCACCAGCG TTGCCAATTG CTGCCCTGTC	200
CCCAGCCCCA ATGGGGGAGT GAGAGAGGCC ACTGCCGGCC GGAC	244
ATG GGT CTC CCC ATC GTG CCT GGC CTG CTG CTG TCA CTG GTG CTC	289
Met Gly Leu Pro Ile Val Pro Gly Leu Leu Ser Leu Val Leu	
5 · 10 15	
CTG GCT CTG CTG ATG GGG ATA CAC CCA TCA GGG GTC ACC GGA CTG	334
Leu Ala Leu Leu Met Gly Ile His Pro Ser Gly Val Thr Gly Leu	

20 25 30 GTT CCT TCT CTT GGT GAC CGG GAG AAG AGG GAT AAT TTG TGT CCC Val Pro Ser Leu Gly Asp Ara Glu Lys Arg Asp Asn Leu Cys Pro 35 45 CAG GGA AAG TAT GCC CAT CCA AAG AAT AAT TCC ATC TGC TGC ACC Gln Gly Lys Tyr Ala His Pro Lys Asn Asn Ser Ile Cys Cvs Thr 50 55 AAG TGC CAC AAA GGA ACC TAC TTG GTG AGT GAC TGT CCA AGC CCA Lys Cys His Lys Gly Thr Tyr Leu Val Ser Asp Cys Pro Ser Pro GGG CAG GAA ACA GTC TGC GAG CTC TCT CAT AAA GGC ACC TTT ACA 514 Gly Gln Glu Thr Val Cys Glu Leu Ser His Lys Gly Thr Phe Thr 80 90 GCT TCG CAG AAC CAC GTC AGA CAG TGT CTC AGT TGC AAG ACA TGT Ala Ser Gln Asn His Val Arg Gln Cys Leu Ser Cys Lys Thr Cys 100 CGG AAA GAA ATG TTC CAG GTG GAG ATT TCT CCT TGC AAA GCT GAC Arg Lys Glu Yet Phe Gln Val Glu Ile Ser Pro Cys Lys Ala Asp 110 115 120 ATG GAC ACC GTG TGT GGC TGC AAG PAG AkC CAA TTC CAG CGC TAC Met Asp Thr Val Cys Gly Cys Lys Lys Asn Gln Phe Gln Arg Tyr 125 130 135 CTG AGT GAG ACG CAT TTC CAG TGT GTG GAC TGC AGC CCC TGC TTC Leu Ser Glu Thr His Phe Gln Cys Val Asp Cys Ser Pro Cys Phe 140 145 150 AAT GGC ACC GTG ACA ATC CCC TGT AAG GAG AAA CAG AAC ACC GTG 739 Asn Gly Thr Val Thr Ile Pro Cys Lys Glu Lys Gln Asn Thr Val 155 160 165 TGT AAC TGC CAC GCA GGA TTC TTT CTA AGC GGA AAT GAG TGC ACC Cys Asn Cys His Ala Gly Phe Phe Leu Ser Gly Asn Glu Cys Thr 175 180 CCT TGC AGC CAC TGC AAG AAA AAT CAG GAA TGT ATG AAG CTG TGC Pro Cys Ser His Cys Lys Lys Asn Gln Glu Cys Met Lys Leu Cys 185 190 195 CTA CCT CCA GTT GCA AAT GTC ACA AAC CCC CAG GAC TCA GGT ACT

Leu Pro	Pro Val	Ala Asn 200	Val Thr	Asn	Pro 205	Gln	Asp	Ser	Gly	Thr 210	
		CCT CTG Pro Leu 215									919
		ATC AGT Ile Ser 230									964
		TCC ATC Ser Ile 245									1009
		GAA GGA Glu Gly 260									1054
		TTC AGC Phe Ser 275									1099
		ACC CCA Thr Pro 290									1144
		GTC TTC Val Phe 305									1189
		GAG GTG Glu Val 320									1234
		CTC AAC Leu Asn 335									1279
		GTC GTC Val Val 350									1324
		ATG CTG Met Leu 365									1369

		TGG Trp												CAC His	1414
		GAG Glu												GAG	1459
GIU	116	Giu	ALC	395	Giu	пец	GIII	ASII	400	Arg	Cys	ьец	Arg	405	
GCT	CAT	TAC	AGC	ATG	CTG	GAA	GCC	TGG	CGG	CGC	CGC	ACA	CCG	CGA	1504
Ala	His	Tyr	Ser	Met 410	Leu	Glu	Ala	Trp	Arg 415	Arg	Arg	Thr	Pro	Arg 420	
an a	a 7 a	aaa	7.00	СШС	a	CITI 3	СШС	aaa	aaa	аша	OIIII	maa	a	3 mg	1 = 40
		Ala													1549
	0_0			425	1101		7	U -1	430			Cyb	p	435	
AAC	CTG	CGT	GGC	TGC	CTG	GAG	AAC	ATC	CGC	GAG	ACT	CTA	GAA	AGC	1594
Asn	Leu	Arg	Gly	_	Leu	Glu	Asn	Ile	_	Glu	Thr	Leu	Glu		
				440					445					450	
CCT	GCC	CAC	TCG	TCC	ACG	ACC	CAC	CTC	CCG	CGA	TAAC	GCCI	ACA		1637
Pro	Ala	His	Ser		Thr	Thr	His	Leu		Arg					
				455					460						
CCC	CCAC	CTC A	AGGA	ACGGC	A CI	CGA	AGGAC	CAT	CCTC	GCTA	GATO	CCC?	rgc		1687
TTC	CCTGT	rga <i>i</i>	ACCTO	CTCT	T TO	GTC	CTCTA	A GGC	GGC <i>I</i>	AGGC	TCGA	ATCTO	GC.		1737
AGG	CTCGA	ATC I	rggc <i>i</i>	AGCC	AC TI	CCTT	GGTG	G CTA	ACCGA	ACTT	GGT	TAC	ATA		1787
GCTT	CTTC	CCA C	CTGC	CCGAC	G AC	CAGC	CTGTC	CCA	AGCC	ACTT	GTG	CATGO	GCA		1837
GGG	AGTO	TG (CCATO	CTGCT	C CC	CAGAC	CAGCT	GAG	GGT	GCCA	AAAC	CCAC	3GA		1887
GAG	GTGA'	TG 1	rggac	AAAA	AA GC	CACA	ATCTA	A TCT	GAT!	ACCC	ACT	rggg <i>i</i>	ATG		1937
CAAC	GACC	CCA A	ACAA	AGCI	T C	CAG	GCC1	CCI	CAG	rtga	TTTC	CTGGC	GCC .		1987
CTTT	CTCAC	CAG 1	[AGA]	'AAAA'	AC AC	TCT	TGT	A TTC	ATTA	TAT	CAC	ACTAZ	ATG		2037
GATO	BAACC	GT 1	rgaac	CTCCC	CT AF	AGGT	AGGGG	CAA	AGCA	CAGA	ACAC	TGGC	GT		2087
CTCC	CAGC	rgg <i>i</i>	AGCCC	CCGA	C TO	CTTGT	CAAA'	ACA	ACTAZ	AAAA	TCT	AAAA	GTG		2137
AAAA	AAAA	AAA A	AAAA	AAAA	AA AA	AAAA	AAAA	GAA	ATTC						2173

(70) INFORMATION FOR SEQ ID NO:69:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 461 amino acids
- (B) TYPE:polypeptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:

Met Gly Leu Pro Ile Val Pro Gly Leu Leu Leu Ser Leu Val Leu 5 10 15 Leu Ala Leu Leu Met Gly Ile His Pro Ser Gly Val Thr Gly Leu

Val Pro Ser Leu Gly Asp Ara Glu Lys Arg Asp Asn Leu Cys Pro Gln Gly Lys Tyr Ala His Pro Lys Asn Asn Ser Ile Cys Cvs Thr Lys Cys His Lys Gly Thr Tyr Leu Val Ser Asp Cys Pro Ser Pro Gly Gln Glu Thr Val Cys Glu Leu Ser His Lys Gly Thr Phe Thr Ala Ser Gln Asn His Val Arg Gln Cys Leu Ser Cys Lys Thr Cys Arg Lys Glu Yet Phe Gln Val Glu Ile Ser Pro Cys Lys Ala Asp Met Asp Thr Val Cys Gly Cys Lys Lys Asn Gln Phe Gln Arg Tyr Leu Ser Glu Thr His Phe Gln Cys Val Asp Cys Ser Pro Cys Phe Asn Gly Thr Val Thr Ile Pro Cys Lys Glu Lys Gln Asn Thr Val Cys Asn Cys His Ala Gly Phe Phe Leu Ser Gly Asn Glu Cys Thr Pro Cys Ser His Cys Lys Lys Asn Gln Glu Cys Met Lys Leu Cys Leu Pro Pro Val Ala Asn Val Thr Asn Pro Gln Asp Ser Gly Thr Ala Val Leu Leu Pro Leu Val Ile Phe Leu Gly Leu Cys Leu Leu Phe Phe Ile Cys Ile Ser Leu Leu Cys Arg Tyr Pro Gln Trp Arg Pro Arg Val Tyr Ser Ile Ile Cys Arg Asp Ser Ala Pro Val Lys Glu Val Glu Gly Glu Gly Ile Val Thr Lys Pro Leu Thr Pro Ala Ser Ile Pro Ala Phe Ser Pro Asn Pro Gly Phe Asn Pro Thr Leu Gly Phe Ser Thr Thr Pro Arg Phe Ser His Pro Val Ser Ser Thr Pro Ile Ser Pro Val Phe Gly Pro Ser Asn Trp His Asn Phe Val Pro Pro Val Arg Glu Val Val Pro Thr Gln Gly Ala Asp Pro Leu Leu Tyr Gly Ser Leu Asn Pro Val Pro Ile Pro Ala Pro Val Arg Lys Trp Glu Asp Val Val Ala Ala Gln Pro Gln Arg Leu Asp Thr Ala Asp Pro Ala Met Leu Tyr Ala Val Val Asp Gly Val Pro Pro

				365					370					375
Thr	Arg	Trp	Lys	Glu	Phe	Met	Arg	Leu	Leu	Gly	Leu	Ser	Glu	His
				380					385					390
Glu	Ile	Glu	Arc	Leu	Glu	Leu	Gln	Asn	Gly	Arg	Cys	Leu	Arg	Glu
				395					400					405
Ala	His	Tyr	Ser	Met	Leu	Glu	Ala	Trp	Arg	Arg	Arg	Thr	Pro	Arg
				410					415					420
His	Glu	Ala	Thr	Leu	Asp	Val	Val	Gly	Arg	Val	Leu	Cys	Asp	Met
				425					430					435
Asn	Leu	Arg	Gly	Cys	Leu	Glu	Asn	Ile	Arg	Glu	Thr	Leu	Glu	Ser
				440					445					450
Pro	Ala	His	Ser	Ser	Thr	Thr	His	Leu	Pro	Arg				
				455					460					

(71) INFORMATION FOR SEQ ID NO:70:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH:1232 bases
- (B) TYPE:nucleic acid
- (C) STRANDEDNESS:single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:

GAA?	TCTC	CTG (GACTO	GAGG(CT CC	CAGT	CTGC	G CCI	rttgo	GGT	TCA	AGAT(CAC		50
TGGC	GACCA	AGG (CCGT	GATC:	C TA	ATGC	CCGAC	TC	rcaa(CCCT	CAA	CTGT	CAC		100
CCCA	AAGG	CAC :	rtggc	GACG:	rc ci	rgga(CAGA	CGA	AGTC	CCGG	GAA	GCCC	CAG		150
CACI	rgcc	GCT (GCCAC	CACTO	GC CC	CTGAC	GCCC	A GAT	rggg	GGAG	TGAG	GAGG	CCA		200
TAGO	CTGT	CTG (G C												212
															257
Met	Gly	Leu	Ser	Thr	Val	Pro	Asp	Leu	Leu	Leu	Pro	Leu	Val	Leu	
				5					10					15	
										•					
			TTG												302
Leu	Glu	Leu	Leu	Val	Gly	Ile	Tyr	Pro	Ser	Gly	Val	Ile	Gly	Leu	
				20					25					30	
			CTA												347
Val	Pro	Hıs	Leu	_	Asp	Arg	GIu	Lys	_	Asp	Ser	Val	Cys		
				35					40					45	
C 3 3	aa	777	ma m	7 m.c	G3.G	a a m	C13.3	3 3 CC	3 3 CD	шаа	3 mm	maa	mam	3.00	200
			TAT												392
GIN	GIY	гуѕ	Tyr		HIS	Pro	Gin	Asn		ser	тте	Cys	Cys		
				50					55					60	
ልልር	тсс	CAC	AAA	CCA	ACC	ጥ ል ር	<u> ምጥር</u>	ጥ ል ር	יי מ מ	CAC	ጥረረጥ	CCA	GGC	CCG	437
* 77.70		\sim	4747	-	$\sim \sim 1$	$\tau \tau \sim$	440	$\perp \sim$	1777	$\Box \Box \Box$	101	-	\sim	\sim	TJ /

Lys	Cys	His	Lys	Gly 65	Thr	Tyr	Leu	Tyr	Asn 70	Asp	Cys	Pro	Gly	Pro 75	
												TCC Ser			482
												TCC Ser			527
												ACA Thr			572
												CGG Arg			617
				125					130					135	
												CTC Leu			662
												AAC Asn			707
												GAG Glu			752
												AAG Lys			797
												TCA Ser			842
												TGC Cys			887
												CGG Arg			932

		CTC Leu										977
		GAG Glu										1022
		TTC Phe										1067
		GTG Val										1112
		GAC Asp										1157
		TAT Tyr										1202
TCC	GAC	CCC	ATC	CCC	AAC	CCC	CTT	CAG	AAG			1232
Ser	Asp	Pro	Ile	Pro	Asn	Pro	Leu	Gln	Lys			
				335					340			

- (72) INFORMATION FOR SEQ ID NO:71:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 340 amino acids
- (B) TYPE:polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:

Met Gly Leu Ser Thr Val Pro Asp Leu Leu Leu Pro Leu Val Leu
5 10 15

Leu Glu Leu Leu Val Gly Ile Tyr Pro Ser Gly Val Ile Gly Leu
20 25 30

Val Pro His Leu Gly Asp Arg Glu Lys Arg Asp Ser Val Cys Pro
35 40 45

Gln Gly Lys Tyr Ile His Pro Gln Asn Asn Ser Ile Cys Cys Thr

Lys Cys His Lys Gly Thr Tyr Leu Tyr Asn Asp Cys Pro Gly Pro Gly Gln Asp Thr Asp Cys Arg Glu Cys Glu Ser Gly Ser Phe Thr Ala Ser Glu Asn His Leu Arg His Cys Leu Ser Cys Ser Lys Cys Arg Lys Glu Met Gly Gln Val Glu Ile Ser Ser Cys Thr Val Asp Arg Asp Thr Val Cys Gly Cys Arg Lys Asn Gln Tyr Arg His Tyr Trp Ser Glu Asn Leu Phe Gln Cys Phe Asn Cys Ser Leu Cys Leu Asn Gly Thr Val His Leu Ser Cys Gln Glu Lys Gln Asn Thr Val Cys Thr Cys His Ala Gly Phe Phe Leu Arg Glu Asn Glu Cys Val Ser Cys Ser Asn Cys Lys Lys Ser Leu Glu Cys Thr Lys Leu Cys Leu Pro Gln Ile Glu Asn Val Lys Gly Thr Glu Asp Ser Gly Thr Thr Val Leu Leu Pro Leu Val Ile Phe Phe Gly Leu Cys Leu Leu Ser Leu Leu Phe Ile Gly Leu Met Tyr Arg Tyr Gln Arg Trp Lys Ser Lys Leu Tyr Ser Ile Val Cys Gly Lys Ser Thr Pro Glu Lys Glu Gly Glu Leu Glu Gly Thr Thr Thr Lys Pro Leu Ala Pro Asn Pro Ser Phe Ser Pro Thr Pro Gly Phe Thr Pro Thr Leu Gly Phe Ser Pro Val Pro Ser Ser Thr Phe Thr Ser Ser Ser Thr Tyr Thr Pro Gly Asp Cys Pro Asn Phe Ala Ala Pro Arg Arg Glu Val Ala Pro Pro Tyr Gln Gly Ala Asp Pro Ile Leu Ala Thr Ala Leu Ala Ser Asp Pro Ile Pro Asn Pro Leu Gln Lys

(73) INFORMATION FOR SEQ ID NO:72:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 19 bases

(B) TYPE:nucleic acid

- (C) STRANDEDNESS:single
- (D) TOPOLOGY:linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:

GTACTTGAAC TCGTTCCTG

19

Appendix D





STATE OF ISRAEL

This is to certify that nanered bereto is a true topy of the documents as riginally deposited with e patent application eticulars of which are cified on the first page the enner.

זאת לתעודה כי רצופים בזה העתקים נבתים של המסמכים שהופקרו לכתחילה עם חבקשה לפטנט לפי הפרטים הרשומים בעמוד הראשון של חנספת



Registrar of Patents

לשימוש הלשבה For Office Use

092697 מספר: Number 1 3 -12- 1989 :תאריך Dele הוקרם/נוחה Ante/Post-deled

... חוק הפטנטים. תשכ"ו – 1967 PATENT LAW, 5727 - 1967

לפסנס

Application for Palent

אני. (שם המבקש. מענר ולנבי נוף מאונד - מקום החאנרותו) I (Name and address of applicant, and in case of body corporate-place of incorporation)

Yeda Research and Development Co. Ltd. A company registered under the laws of Israel

P,O. Box 95, Rehovot, Israel

ידע חברה למחקר ופיתוח בע"מ. חברה רשומה בישראל 95 .T.n

Assignment by the inventors of an invention the title of which is

העברה מהממציאים

בעל אמבאה מכח... Owner, by virtue of

רחובות

שיבוט מולקולרי של חלבון הקושר קאד (בעברית) (Hobrow)

> (באנגליה) (English)

Molecular cloning of TNP Binding Protein

Application of Division Application for Patent Addition	;	את דין קדימה Priority Cla	
from Application to Patent/Appl.	18°8/1600 Number/Merk	קריך Date	Convention Country
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י פור כו: כללי / מיחד – רבוף בזה / עוד יונש P.O.A.: general/individual-attached/to be filed later- הובש בענין			
Address for Service in brook aulina Ben-Ami, Patent Attorney, iter-Lab Ltd., Kiryat Weizmann,			
ess-Ziona 76110, Israel	j		
Signature of Applicant For the Applicant: ()		989 אנבר F of the yesDece	בחורש דענ
and hybridants y		or the yearect	ember of 11 לשימים הלשכה For Office Use
Paulina Ben-Ami			לשימוש הלשכה

שיבוט מולקולרי של חלבון הקושר TNF Binding Protein

Yeda Research and Development Co. Ltd. T/811 The invention relates to Tumor Necrosis Pactor (TNF) Binding Protein I and more particularly, to the cloning of gene coding for said protein.

Patent Application No. 83878 of the same applicant discloses a new protein found in urine and capable of inhibiting the binding of TNF to its receptors and the cytotoxic effect of TNF. This protein is now referred to hereinafter as TNF Binding Protein I or TBP-I.

The process for the extraction and purification of TBP-I in the above mentioned patent application comprises the following steps:

- (a) recovering the crude protein fraction from a dyalized concentrate of human urine;
- (b) subjecting said crude protein fraction of step (a) to ion exchange chromatography to obtain partially purified active fractions of the TNF Binding Protein defined by its ability to inhibit both the binding of TNF to its receptors and the cytotoxic effect of TNF:
- (c) applying said partially purified active fractions of the TNP Binding Protein from step (b) to reversed phase high pressure liquid chromatography (HPLC) to obtain substantially purified active fractions of the TNP Binding Protein defined by its ability to inhibit both the binding of TNP to its receptors and the cytotoxic effect of TNP; and
- (d) recovering the substantially purified protein of step (c), said protein having a molecular weight of about 26-28 Kda on SDS PAGE under reducing conditions, moving as a single peak on reversed phase HPLC and having the ability to inhibit both the binding of TNF to its receptors and the cytotoxic effect of TNF.

The purified TBP-I was sequenced and shown to contain at the N-terminus the following amino acid sequence:

1 5 10 15
Asp-Ser-Val-Cys-Pro-Gln-Gly-Lys-Tyr-Ile-His-Pro-Gln-Asn-Asn-Ser

The invention relates to oligonucleotide probes to the cDNA coding for a protein comprising the amino acid sequence of TBP-1. The probes were synthesized by known methods on the basis of the above amino acid sequence of the N-terminus of TBP-I.

The invention also relates to a DNA molecule comprising a recombinant DNA molecule or a cDNA molecule coding for a protein comprising the amino acid sequence of TBP-I or a protein substantially homologous therewith. Within the scope of the invention are DNA molecules encoding said homologous proteins having the same biological activities of TBP-I

In a preferred embodiment, the DNA molecule is a cDNA molecule picked up from a human cDNA library, in particular a colon cDNA library. Illustrated in Figure 1 is a partial restriction map of an insert of about 1.0 Kb of such a cDNA molecule obtained in agarose gel and herein designated C2. Figure 2 illustrates a partial nucleotide sequence of said C2 insert and also a partial translated amino acid sequence comprising the NH₂-terminal amino acid sequence of TBP-I encoded thereby. Figure 3 illustrates another partial nucleotide sequence of the C2 insert starting from nucleotide 342. Figure 4 shows a possible nucleotide sequence of the whole insert, that seems to have 965 nucleotides.

The invention further comprises cloning of said cDNA molecule

into a replicable plasmid vector and transformation of a bacterium, e.g., competent E.coli TG1 therewith.

In another aspect, the invention comprises the isolation of mRNA coding for a protein comprising the amino acid sequence of TBP-I by extraction from cells and its detection by hybridization with the cDNA of the invention.

Once the mRNA is obtained in a purified form, the cDNA coding for a protein comprising the amino acid sequence of TBP-I can be obtained by contacting the mRNA with reverse transcriptase for a time and under conditions sufficient to form said cDNA. This cDNA may be converted to double stranded cDNA by known techniques.

Probes may be prepared from the cDNA sequences of the invention and used for isolation of the genomic DNA coding for a protein comprising the amino acid sequence of TBP-I by known methods, e.g. by colony hybridization techniques under stringent conditions.

The DNA of positive clones are then inserted into appropriately constructed expression vectors by techniques well known in the art. Double-stranded cDNA is linked to plasmid vectors by homopolymeric tailing or by restriction linking involving the use of synthetic DNA linkers or blunt-ended ligation techniques. DNA ligases are used to ligate the DNA molecules and undesirable joining is avoided by treatment with alkaline phosphatase.

In order to be capable of expressing a desired protein, an expression vector should comprise also specific nucleotide sequences containing transcriptional and translational regulatory information linked to the DNA coding for the desired protein in such a way as to permit gene expression and production of the protein. The gene must be preceded by a promoter in order to be transcribed. There are a

variety of such promoters in use, which work with different efficiencies (strong and weak promoters).

The DNA molecule comprising the nucleotide sequence coding for a protein comprising the amino acid sequence of TBP-I preceded by a nucleotide sequence of a signal peptide and the operably linked transcriptional and translational regulatory signals is inserted into a vector which is capable of integrating the desired gene sequences into the host cell chromosome. The cells which have stably integrated the introduced DNA into their chromosomes can be selected by also introducing one or more markers which allow for selection of host cells which contain the expression vector.

In a preferred embodiment, the introduced DNA molecule will be incorporated into a plasmid or viral vector capable of autonomous replication in the recipient host. Factors of importance in selecting a particular plasmid or viral vector include the ease with which recipient cells that contain the vector may be recognized and selected from those recipient cells which do not contain the vector; the number of copies of the vector which are desired in a particular host and whether it is desirable to be able to "shuttle" the vector between host cells of different species. Once the vector or DNA sequence containing the construct(s) has been prepared for expression, the DNA construct(s) may be introduced into an appropriate host cell by any of a variety of suitable means: transformation, transfection, conjugation, protoplast fusion, electroporation, calcium phosphate precipitation, direct microinjection, etc.

Host cells to be used in this invention may be either prokaryotic or eukaryotic. Preferred prokaryotic hosts include bacteria, such as E.coli. Under such conditions, the protein will

not be glycosylated. The prokaryotic host must be compatible with the replicon and control sequences in the expression plasmid.

Preferred eukaryotic hosts are mammalian cells, e.g., human, monkey, mouse and chinese hamster ovary (CHO) cells, because they provide post-translational modifications to protein molecules including correct folding or glycosylation at correct sites. Also yeast and insect cells can carry out post-translational peptide modifications including glycosylation. A number of recombinant DNA strategies exist which utilize strong promoter sequences and high copy number of plasmids which can be utilized for production of the desired proteins in yeast. Yeast recognizes leader sequences on cloned mammalian gene products and secretes peptides bearing leader sequences (i.e. pre-peptides).

After the introduction of the vector, the host cells are grown in a selective medium, which selects for the growth of vector-containing cells. Expression of the cloned gene sequence(s) results in the production of the desired protein or a fragment thereof. The expressed protein is then isolated and purified by any conventional procedure involving extraction, precipitation, chromatography, electrophoresis, or the like.

The invention will be illustrated by the following examples:

Example 1

Preparation of oligonucleotide probes

Oligonucleotide probes to the cDNA of TBP-I were designed on the basis of the NH₂-terminal amino acid sequence of the protein. Three mixtures of the synthetic oligonucleotides shown below and designated 1008, 1009, 1010 were used.

Probe 1008 is a mixture of 64 different 26-mers, in which deoxyinosine was introduced wherever the number of possible alternative codons for the amino acid exceeded 3. The two other mixtures are 17-mers; the first, probe 1009, is a mixture of 128 different oligonucleotides and the second, probe 1010, of 196 different oligonucleotides. Each of these two latter mixtures corresponds to part of the amino acid sequence coded-for by 1008. The nucleotide sequence of these two mixtures overlap each other.

GGI GTC CCI TTC ATA TAA GTA GGI GT
T T G G G
T

1009

GGA GTC CCA TTC ATA TA
C T C T G
T G G C
T T G G G C
T G
T G G G C
T G

Example 2

Isolation of cDMA clones

cDNA clones comprising a nucleotide sequence coding for a protein comprising the amino acid sequence of TBP-I were isolated from a human cDNA colon library with the aid of the oligonucleotide probes of Example I as follows:

Four cDNA libraries constructed in lambda gtll (Clontech Laboratories, Inc., U.S.A.) derived from the mRNA of human liver, human placenta, human colon and of HeLa cells were screened with the aid of the 1008 probes of Example 1. The liver, placenta and HeLa cDNA libraries were oligo dT primed, while the colon cDNA library was randomly primed. In each screening, 5x10° phages were adsorbed to

W

Escherichia coli, strain Y1088, plated at a density of 40,000 p.f.u/15 cm petri dish and grown at 37°C for 18 hours. Nitrocellulose filters were overlaid in duplicates on the plates, then immersed in DNA-denaturing solution, transferred further to a neutralizing solution, and then dried in vacuum at 80°C and prehybridized to allow non-specific sites to be saturated with unlabelled DNA. The 1008 probes were 32P-end-labelled, using the T4 polynucleotide kinase and applied to the filters in a solution containing 6 SSC (1 SSC corresponds to 0.15M NaCl and 0.015M sodium citrate), 10 *Denhardt*s solution* (a mixture of Ficoll, polyvinylpyrrolidone and bovine serum albumin (Pentax Fraction V) in water, according to T. Maniatis, et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press, N.Y., 1982 p. 448), 0.5% SDS (sodium dodecyl sulfate), and 100 μg/ml salmon sperm DNA. Hybridization was carried out for 18 hours at 50°C with the colon library and at 42°C with the other libraries. Unbound labelled probe was washed with a solution containing 3SSC at 25°C and then twice again either at 42°C or, for the colon library, at 50°C. Positive clones, identified by exposure of the filters to autoradiography, were picked purified up, and checked hybridization to the 1009 and 1010 probes under those same conditions which were applied for the screening with the 1008 probes, except that the temperature of hybridization and washing was 30°C.

The results of the screening are summarized in Table I. Clones which hybridized with all three probes could be detected only in the placenta and colon libraries. In further analysis of the nucleotide sequence, only the clones picked up from the colon library (the only library which was randomly primed) were found indeed to code for TBP-I. These clones were designated C1, C2, C3, and C4.

Table I: Libraries screened for the TBP-I cDNA and the clones which were isolated

Vector	Library	Clone na	ime Temp. of 1008 C°	hybridiza 1009 C°	tion 1010 C°
Agtll cDNA oligo dT primed		į			
	Liver	-	-	-	-
	HeLa	-	-	-	-
	Placenta	17 19 131 133 152	50° 50° 60° 50°	30° 30° 30°	30° 30°
Agt11 cDNA randomly primed					-
normal tissue arou	Colon	C1 C2 C3 C4	60° 60° 60°	30°	30°. 30° 30°

Example 3

Characterization of the isolated clones from the human colon cDNA library

The purified lambda gtll DNA containing positive cDNA clones were digested with EcoRl and size-fractionated on 11 agarose gel. Two of the clones had an insert size of about 1.0 Kb, the third was of 0.9 Kb and the fourth had two inserts of 0.9 and 0.8 Kb. Cross-hybridization among the four clones was tested by Southern blotting. The results are summarized in Table II. Clones C2 and C3 were found to contain the same 1.0 Kb insert. The restriction map of this insert is shown in Figure 1. Clone C5 contains two inserts, of 0.9 and 0.8 Kb: the 0.9 Kb insert constitutes part of the insert of C2, while the 0.8 Kb insert seems to be unrelated. Clone C1 also contains

a 0.9 Kb insert which constitutes part of the insert of C2.

Table II: Insert sizes and interrelationships in the various cDNA clones for TBP-I

cDNA clone	Insert size (Kb)	Cr with the Cl insert	oss hybridiz with the C2 insert	ation: - with the EcoRI PetI 165 nucleotide - fragment of C2
C1	0.9	++	+	
C2	1.0	+	++	++
C3	1.0	+	++	++
C5	0.9, 0.8	+	++	++

The 1.0 Kb EcoRI insert of the C2 clone was subcloned in a Bluescript plasmid vector of Stratagene Cloning System (San Diego, Cal.) and E. coli TG1 competent bacteria were transformed therewith. The transformed bacteria were deposited with the Collection Nationale de Cultures de Microorganismes (C.N.C.M.), of the Institut Pasteur, Paris, France, on December 6, 1989 under the Budapest Treaty, and it was assigned the deposit number CNCM I-917.

Example 4

. Screening of an oligo dT primed (human placents) cDNA library with the DNA probes

To isolate sequences extending 3' to the insert of clone C2,

0.5x10° recombinants from human placenta cDNA library in lambda gtll

were screened for hybridization with the aid of a probe prepared from

the 125-nucleotide PstI-PstI fragment of clone C2 (see Fig. 1) which

was labelled with the multiprime DNA labelling systems kit

(Amersham). The technique is based on the use of random sequence hexanucleotide to prime DNA synthesis on denatured template DNA at numerous sites along its length.

Phages were adsorbed to E.coli strain Y1088, plated at a density of 40,000 p.f.u/15 cm petri dish and grown at 37°C overnight. Two sets of nitrocellulose filters were overlaid and immersed in a tray containing DNA-denaturing solution. The filters were washed, fixed, neutralized, dried at 80°C under vacuum and prehybridized to allow non-specific sites to be saturated by unlabelled DNA. Then the filters were hybridized with the ***P-labelled probe overnight at 65°C. Unbound label was washed first in a solution containing ISSC and 0.1% SDS (twice at 25°C and then twice again at 65°C) and then . at 65°C in a solution containing 0.1 SSC and 0.1% SDS. Filters were autoradiographed. Thirteen positive clones were obtained and picked up. After purification, these clones were tested for hybridization with a probe constructed from the C2 insert from which the above EcoRI-PstI insert was deleted. Four clones were found to hybridize. DNA from these positive clones was isolated after purifying the phages by centrifugation in a CsCl solution. Their inserts were excised by cutting with EcoRI and their sizes were estimated by electrophoresis on 12 agarose gel. The phage containing the largest fragment of about 2.0 Kb was subjected to further analysis. Its insert was subcloned in a Bluescript plasmid vector and E.coli TG 1 competent bacteria were transformed therewith.

Example 5

Determination of the nucleotide sequence in the cloned DNA

DNA of the 1.0 Kb EcoRI insert of the C2 clone propagated in the

3

Bluescript plasmid vector was subjected to partial degradation to various extents using the Erase-a-Base Progema kit. DNA of plasmids containing the insert at various degrees of degradation was denatured and subjected to sequencing as described by Hattoni and Sakaki (Anal. Biochem. 152, 232-238) with the aid of the Sequenase Kit (USB).

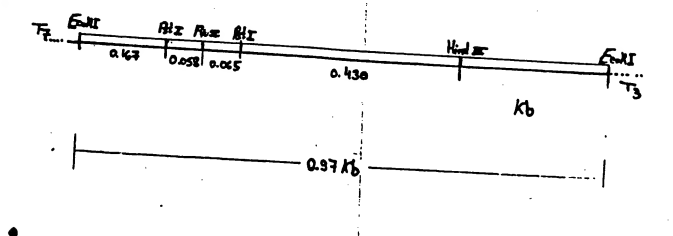
The insert was shown to have 965 nucleotides. Figure 2 shows a partial nucleotide sequence of the C2 insert and the deduced amino acid sequence which it codes for. Within this sequence the NH₂-terminal amino acid sequence of TBP-I, on the basis of which the cDNA had been cloned, can be detected (underlined). Figure 3 shows the 343-965 nucleotide sequence of the C2 insert and Figure 4 shows a possible total nucleotide sequence of the C2 insert.

Example 6

Detection and sizing of the mRNA for TBP-I by Northern blot analysis

Total RNA was extracted from cells of the U937 and HT29 lines by the "hot phenol" method according to T. Maniatis et al., op. cit., p. 194. Samples of 50 and 25µg RNA were analyzed by electrophoresis on 1.51 agarose gel in the presence of 2.21 formaldehyde followed by blotting to charged nylon filter. The EcoRI insert of the C2 clone was 32P-labeled with the use of the multiprime DNA labelling systems kits and hybridized to the charged nylon blot (42°C in 502 formamide). As shown in Figure 5 in both cells the cDNA was found to hybridize to mRNA of just a single size, of about 195-21S and having about 2500 nucleotides (right stronger dot - 50 µg RNA, left weaker dot - 25 µg RNA).

FIGURE 1



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					•
				GG TCAAT GCAGGGTCT	
	•- - .		į.	COMAGTIMACGTCGCACA	360
	;	361	GCCTCAATGGGACCGTGCACCTCTCCTGCCAGGA	AlaSerII PAlaAlage	_
		• • •	COGAGTTACCCTGGGACGTGGAGAGGACGGTCGT AlaserHetGlyProCysThrScrProAleAnna	SAAACAGAASAGGTC 19CACCTGCC	• •
			AlaSerMetGlyProCysTheCong	CTTTGTCTTGTGCCACCCCTGCACCC	3113
		121	ATGCGATTCTTCTCAGAGAAAACCAATT	rgasnarglh-ProCysAlaProAls -	
	-	. 2 1	TACGCTAAGAAAGATTCTCTTTTGCTCACACACACACACA	CTGTAGTAAT TSTAGGAAAAGCCTA	`. :
			TACGCTAAGAAAGATTCTCTTTTGCTCACAGAGAGAGAGA	GACATCATIGALATTCTTTTCGGGC AL	ыņ
			HetargPhePheLeuargGluAsnGluCysValSa GAGTGCACGAAGTTGTGCCTACCCCAGATTGACA	Programment of the Seel of the	
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			GTCACGTGCTTCAACACGGATGGGGTCTAACTCTT	ACANTTCCCGTGACTUC	10
			GlucysThrLysLauCysLauProGluIleGluAs ACCACAGTCGTGTTCAAAATGGTCATTTTCTTTC	NValimenturian	
	54	41	ACCACAGTCGTGTTCAAAATGGTCATTTTCTTTGU TGGTGTCAGCACAACTTTTACCAGTAAAAAAAAA	TCT:TCC:::	
		_	TGGTGTCAGCACAACTTTTACCAGTAAAAGAAACC	ACADA	· .
			The The Valuable of Land Control of the Control of	AGAAACGRAAATAUGGAGAGAAG . 60); •
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		Ť	TTTAGCTGTGACTITTCTCCCCTCC	ACTACTARCCCCT DOCCCCAAACC	
		L	TTTAGCTGTGACTITTCTCCCCTCGAACTTCUTTGA	TGATGATTUGGGAUCGGGGTTTCG >>>'C	j
	721	C	LysSerThrLeuLysArgGlyGluLeuGluGlyThr CAAGCTTCAGTCCCACTCCAGGCTTCACCCCCACGC	ThrThrLy-ProlruProGlation	
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		G	PTTCGAAGTCAGGGTGAGGTCCGAAGTGGGGGTGGG	ACCCGNAGT CAGGGCACGCCTC 780	
		Co	CACCTTCACCTCCACCTCCACCTATACCCCCCCCCCCC	rpAleSerValDrocupper	
	781	GG	GTGGAAGTGGAGGTCGAGGTGATATCCCCCGGTGA	CIGICCGAATTICTOCTOCT	
	·	P =	GTGGAAGTGGAGGTCGAGGTGGATATGGGGGCCACT	GACAGGGTTAAAA	
		CA	rgProSerProProAlaProProIlaProProValT AGAGAGGTGGCACCACCTATCAGGGGGCTAACCAC	hrvale attack	
	841		AGAGAGOTGGCACCACCTATCAGGGGGCTGACCCCA TCTCTCCACCGTGGTGGATAGTCCCCCAACTCAGG	TCC Toposous -	
		61	TCTCTCCACCGTGGTGGATAGTCCCCGACTGCGGT	AGGAACGE TOTCO	
		GI	InArgGlyGlyThrThrTyrGlnGlyAlaAcaBack	Tallanda TGTCOGGAGCGGAGA	
9	701	GA	InArgGlyGlyThrThrTyrGlnGlyAlaAspProl ACCCCATCCCAACCCCCTTCAGAAGTGCCACCACA	trieuAlaThrAJul euAlaSur -	
		CTO	GGGGTAGGGGTTGGGGGAAGTCTTCAGG		
		Asp	PProlleProAsnProLaugial war-	ECGGTGTTCGGTGTCTCGGGGC 960	
q	61	AAT	TTC TTC	: PALETATE : HTSETHISTE	
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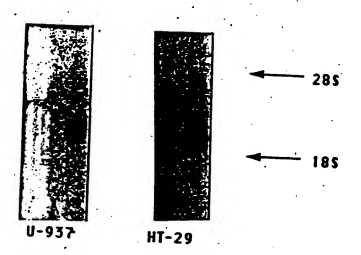
FIGURE 4

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TATEGAATTCCGGTCCCTCACCTAGGGGACAGGGACAGAGAGAG
ATACCTTA ASSESSMENT TO THE TOTAL TOT
HAGGGGAGGGAGTGGATGGCTGTCGCTCTTCTC
ATAGCTTAAGGCCAGGGAGTGGATCCCCTGTCCCCTCTCTCT
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GGAAAATATATCCACCTCAAAATAATTCGAFTTGCTGTACCAAGTGCCAGAAAGGAACC
GLULUSTURIALIS
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121 FIGTIGIACATEACTE TO CARROLLE TO STORY STATE OF STATE
AAGAACATGTTACTGACAGGTCCGGGGCCGCTCCTATGCCTGACGCGACTGTCAGACC Pheleutyraanaspcysproglyproglyctpacate
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THE THE TYPE AS IN A SPECY SPE
PheleuryrAsnAspCysProGlyProGlyGlnAspThrAspCysArgGluCysGluSir -
GGCTCCTTCACGCTICAGAAACCACCTCAGACACTGCCTCAGCTGCTCCCCAATGCCGAAA
CCGAGGAAGTGCGAAGTCTTIGGTGGAGTCTGIGACGGGTCCAGATGCCGAAA GlySyrFhaThrLauglnLysProProGlyThrlaugl
GlySerFhaThrLeuglnLyaProProGluThrLeuplogluLeuLougluMetProLys
GGAATGGTCAGTGAGAGTCTTCTTGCACAGTGGACCGGGACACGTTGTGTGCTGCAGGA
TACCAGTCACTCTGAGAAGAACGTGTCACCTTTTTTTTTT
Grand traiser Grant and and and analysis of the training ACACACACACACACACACACACACACACACACACACAC
GO1 AGAACCAGTACCGGCATTATTGGAGTGAAAACGTTTTCCAGTGGTCAATIGCAGCCTCT
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TCTTGGTCATGGCCGTAATAACCTCACTTTTCGAAAAGGTCACGAAGTTAACGTCGGAGA ArgThrSerThrGly11e11eGlyValLusThrBhag
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ArgThrSerThrGlyIleIleGlyValLyaThrPheSarSerAlaSarIlrAlaAlaSer -
GCCTCAATGGGACCGTGCACCTCTCCTGCCAGGAGAACAGAACAGGACACGTCTGCACCTGCC
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AlaSerMetGlyProCysThrSerProAlaArgArgAsnarglhrProCysAlaProAla
ATGCGATTCTTCTAAGAGAAACGACTGTCTCCCGGTAGTAACTGTAAGAAAAGCCCTG
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GAGTACAGA
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CTCACGTGCTTCAACACGGATGGGGTCTAACTCTTACAATICCCGTCACTCCIGAGTCUG
THACTOTTACANTICCCSTCACTCCIGAGTOCC

	Glucustes
	STORY TOTAL SECUCION OF THE SE
5	GlucyeThrLysLeuCysLeuProGinIleGluAenValLyeGluThrGluAspSerGly -
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60	ATTROTTTAATGTATCGCTACCACCOTOCIACTA
_	TAACCAAATTACATAGCGATGGTTGCCACGTTCCAAGCTCTACTCCATTGTTTGT
	IlegiyLeuMe trurarelurgina-uran
66	AAA1CGACACTGAAAAGAGGGGGGGCTTGAAGCAACTACTACTAACCCCTGACCCCCAAACC
	TTTAGGTGTGAGTTTTGTGGGGTGGAGTTGGTTGATGATG
	LysserThrLauLysArgGlyGluLeuGluGlyThrThrThrLysProlepProGlnThr -
72	CAAGCTTCAGTCCCACTCCAGGCTTCACCCCCCCCCCCC
	GTTCGAAGTCAGGGTGAGGTCCGAAGTGGGGGTGGGACCCGAAGTCAGGCCCAGTT GINALaSerValFroLeugInAlaSerBroRein
	GinAlaSerValFroLeuGinAlaSerProPruProTrpAlaGerValProCysProVal -
781	GCACCTTCACCTCCAGCTCCACCTATACCCCCGGTGACTGTCCCAATTTCTGGCTCCCGG
841	
	CAGAGAGGTGGCACCACCTATCAGGGGGCTGACCCCATCCTTCCGACCAGCCCTCCCCTCC GTCTCTCCACCGTGGTGGATAGTCCCCCGACTGCGGTAGGAACCCTGTCGGGAGCGGACG G1nArgGlyGlyThrThrTyrGlgGlyAl-A
	GINATGGIGGIGTHTTHTTYTGINGIGAIGAGFTAGGAACGCTGTCGGGAGCGGAGG GACCCCATCCCCAACCCCCTTCAGAACTAACAACTAACAACTAACAACTAACAACTAACAACTAACAAC
701	GACCCCATCCCCAACCCCCTTCAGAAGTGGGAGGACACCGCCAGAAGCCACAAGCCACAAGCCACAAGCCACAAGCCACAAGCCACAAGCCACAAGCCACAAGCCACAGCCACAGCCACAAGCCACAGCCACAGAAGCCACAGCCACAGAAGCCACAGCCACAGAAGCCACAGCCACAGAAGCCACAGCCACAGAAGCCACAGCCACAGAAGCCACAGCCACAGAAGA
•	CTGGGGTAGGGGTTGGGGGAAGTCTTCACCCTCCTGTCGCGGTGTTCGGTGTCTCGGCCC
	AspProlleProAsnProLeuGInLyeTrpGJuAspScrAlclhrSerHi:ArgAlaArg -
61	
	TTAAG
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Yeda Research and Development Co. Ltd. T/811

Detection of TBPI mRNA by northern blot hybridization



Yeda Research and Development Co. Ltd. T/811

Claims

- A DNA molecule comprising a recombinant DNA molecule or a cDNA molecule coding for a protein comprising the amino acid sequence of TBP-I or a protein substantially homologous therewith.
- 2. A cDNA molecule according to claim 1 comprising a 1.08 Kb C2 insert of about 0.965 Kb, said insert providing a partial restriction map substantially as shown in Figure 1 upon digestion with restriction enzymes.
- 3. A cDNA molecule according to claim 1 or 2 comprising the partial nucleotide sequence shown in Figure 2 or a nucleotide sequence substantially homologous therewith.
- 4. A cDNA molecule according to claim 1 or 2 comprising the partial nucleotide sequence shown in Figure 3 or a nucleotide sequence substantially homogolous therewith.
- 5. A cDNA molecule according to claim 1 or 2 comprising the partial nucleotide sequences shown in Figures 2 and 3 or nucleotide sequences substantially homogolous therewith.
- 6. A cDNA molecule according to any of the preceding claims comprising the nucleotide sequence shown in Figure 4 or a nucleotide sequence substantially homologous therewith.
- 7. A cDNA molecule according to any of the preceding claims

encoding a protein comprising the amino acid sequence of TBP-I or a protein substantially homologous therewith.

- 8 A replicable plasmid vector comprising a DNA molecule according to any of the preceding claims.
- 9. A replicable plasmid vector according to claim 8 comprising the C2 insert of any of claims 2 to 8.
- 10. A bacterium transformed with a replicable plasmid vector according to claim 8 or 9.
- 11. A bacterium according to claim 10 which is an E.coli strain.
- 12. E.coli TG1 C2 having the deposit number CNCM I-917.
- 13. Oligonucleotide probes useful for picking up genes from cDNA libraries which code for proteins comprising the amino acid sequence of TBP-I, said probes having the formula:

GGI GTC CCI TTC ATA TAA GTA GGI GT
T T G G G
T

1009

GGA GTC CCA TTC ATA TA
C T C T G
T G
G T

1010

TTC ATA TAA GTA GGA GT
T G G G C

14. DNA molecules hybridizable to all three oligonucleotide probes according to claim 13 and which code for a protein comprising

G

the amino acid sequence of TBP-I or a protein substantially homologous therewith.

15. mRNA isolated from U937 and HT29 cells having a size of about 2.5 Kb and being hybridizable with the C2 insert of claims 2 to 7.

16. The mRNA of claim 15 coding for a protein comprising the amino acid sequence of TBP-I or a protein substantially homologous therewith.

For the Applicants:

Paulina Ben-Ami

Patene Attorney

Appendix E

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Soluble forms of tumor necrosis factor receptors (TNF-Rs). The cDNA for the type I TNF-R, cloned using amino acid sequence data of its soluble form, encodes both the cell surface and a soluble form of the receptor

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Communicated by M.Revel

Two proteins which specifically bind tumor necrosis factor (TNF) have recently been isolated from human urine in our laboratory. The two proteins cross-react immunologically with two species of cell surface TNF receptors (TNF-R). Antibodies against one of the two TNF binding proteins (TBPI) were found to have effects characteristic of TNF, including stimulating phosphorylation of specific cellular proteins. Oligonucleotide probes designed on the basis of the NH2-terminal amino acid sequence of TBPI were used to clone the cDNA for the structurally related cell surface type 1 TNF-R. It is notable that although this receptor can signal the phosphorylation of cellular proteins, it appears from its amino acid sequence to be devoid of intrinsic protein kinase activity. The extracellular domain of the receptor is composed of four internal cysteine-rich repeats, homologous to structures repeated four times in the extracellular domains of the nerve growth factor receptor and the B lymphocyte surface antigen CDw40. The amino acid composition and size of the extracellular domain of the type I TNF-R closely resemble those of TBPI. The COOH-terminal amino acid sequence of the four cysteine rich repeats within the extracellular domain of the type I TNF-R matches the COOH-terminal sequence of TBPI. Amino acid sequences in the extracellular domain also fully match other sequences found in TBPI. On the other hand, amino acid sequences in the soluble form of the type II TNF-R (TBPII), while indicating a marked homology of structure, did not suggest any identity between this protein and the extracellular domain of the type I TNF-R. CHO cells transfected with type I TNF-R cDNA produced both cell surface and soluble forms of the receptor. The receptor produced by CHO ceils was recognized by several monoclonal antibodies against TBPI, reacting with several distinct epitopes in this molecule. These data suggest that the soluble forms of the TNF-Rs are structurally identical to the extracellular cytokine binding domains of these receptors and are consistent with the notion that the soluble forms are, at least partly, derived from the same transcripts that encode the cell surface receptors.

Key words: CDw40 antigen/cytokines/nerve growth factor/receptors/tumor necrosis factor

Introduction

Tumor necrosis factors TNF- α and TNF- β (lymphotoxin) are structurally related polypeptide cytokines, produced primarily by mononuclear leukocytes, whose effects on cell function constitute a major factor in the elicitation of the inflammatory response. The TNFs affect cells in different ways, some of which resemble the functional modes of other inflammatory mediators, like interleukin 1 (IL-1) and interleukin 6 (IL-6). What appears most distinctive regarding the activity of the TNFs is that many of their effects can result in cell and tissue destruction. Increasing evidence that over-induction of these destructive activities contributes to the pathogenesis of a number of diseases makes it of particular interest to elucidate their mechanisms and the ways they are regulated (Beutler and Cerami, 1988; Old, 1988). High affinity receptors, to which both TNF- α and TNF- β bind (Baglioni et al., 1985; Beutler et al., 1985; Kull et al., 1985; Tsujimoto et al., 1985; Aggarwal et al., 1986; Israel et al., 1986) play a key role in the initiation and control of the cellular response to these cytokines. These receptors are expressed on the surfaces of a variety of different cells. Studies showing that antibodies reacting with their extracellular portions affect cells in a manner very similar to the TNFs demonstrate that the receptors and cellular components associated with them are sufficient to provide the intracellular signalling for the effects of the TNFs (Engelmann et al., 1990a; Espevik et al., 1990). Other studies have shown that molecules related to the TNF receptors (TNF-Rs) also exist in soluble forms. Two immunologically distinct species of such soluble TNF-Rs, TBPI and TBPII, were recently isolated from human urine (Engelmann et al., 1989, 1990b; Olsson et al., 1989; Seckinger et al., 1989a). Immunological evidence indicated that the two proteins are structurally related to two molecular species of the cell surface TNF-R (the type I and type II receptors, respectively). Antibodies to each of the two soluble proteins were shown to block specifically the binding of TNF to one of the two receptors and could be used to immunoprecipitate the receptors. Antibodies against one of the two soluble proteins (TBPI) were also found to induce effects characteristic of TNF in cells which express the immunologically cross-reactive cell receptors (Engelmann et al., 1990a. 1990b). Like the cell surface receptors for TNF, the soluble forms of these receptors specifically bind TNF and can thus interfere with its binding to cells. It was suggested that they function as physiological inhibitors of TNF activity (Engelmann et al., 1989; Olsson et al., 1989; Seckinger et al., 1989a).

In the present study we explored the structural relationship of the soluble and cell surface forms of the TNF-Rs further by determining amino acid sequences of the soluble forms and by using amino acid sequence data for one of the soluble receptors to clone the cDNA which encodes this protein. Initial information on the mechanism of formation

Y.Nuphar et al.

of the soluble receptors was gained by examining the expression of this cDNA in transfected CHO cells.

Results

Cloning of the cDNA for the type I TNF-R

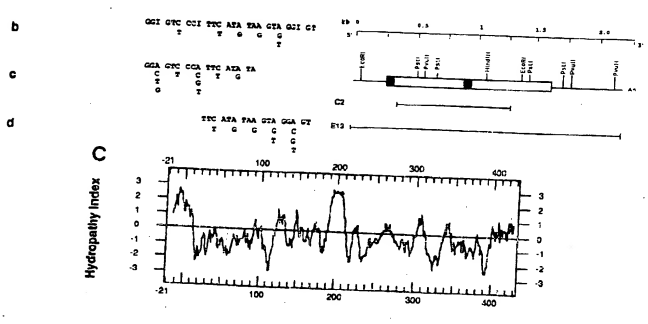
To clone the cDNAs which code for the TNF-binding protein, TBPI, and its related TNF receptor, we screened several cDNA libraries, using three overlapping oligonucleotide probes designed on the basis of the NH2-terminal amino acid sequence of TBPI (Figure 1A). In a \(\lambda GT11 \) library derived from the mRNA of human colon (randomly primed, Clontech, Palo Alto, CA), we detected four recoinbinant phages which hybridized with the three probes. The inserts in these four phages were similar in size, and were found to overlap by restriction mapping and sequence analysis. Complete analysis of the sequence of the longest of the four (C2 in Figure 1B) revealed an open reading frame extending over its entire length. A polypeptide chain encoded in this reading frame fully matches the NH2-terminal amino acid sequence of TBPI. Neither an initiation nor a stop codon was found in the C2 insert. Rescreening the colon cDNA library using another probe corresponding to a sequence found in C2 (see Materials and methods) yielded several other recombinant phages containing inserts that overlap with the C2 insert. However, none of them provided further sequence information on the cDNA in the 5' or the 3' directions. In a AZAP cDNA library derived from the mRNA of CEM lymphocytes (Foley et al., 1965) [oligo (dT) and randomly primed. Clontech] five phages hybridizing with this probe were detected, which contained significantly

longer inserts than C2. The longest insert (E13, Figure 1B) was sequenced in its entirety (Figure 1D) and was found to contain the C2 sequence (nucleotides 346-1277 in Figure 1D) within one long open reading frame of 1365 bp, flanked by untranslated regions of 255 and 556 nucleotides at its 5' and 3' ends respectively. The potential ATG initiation site, occurring at positions 256-258 in the nucleotide sequence. (denoted by an asterisk in Figure 1d) is preceded by an upstream in-frame termination codon at bases 244-246. The start location is in conformity with one of the possible alternatives for the translation initiation consensus sequence (GGCATGG, nucleotides 253-259; Kozak, 1987).

There is no characteristic poly(A) addition signal near the 3' end of the cDNA. The sequence ACTAAA, at nucleotides 2045-2050, may serve as an alternative to this signal, but with low efficiency (Sheets and Wickens, 1989). At nucleotides 1965-2000, there are six consecutive repeats of the sequence G(T)n (n varying between 4 and 8). Similar sequences have also been observed in the 3' noncoding regions of the cDNAs of some members of the jun family, which are also devoid of the characteristic poly(A) signal (Ryder et al., 1988, 1989). The 3' end has a 15 base poly(A)

The size of the protein encoded by the cDNA ($\sim 50 \text{ kd}$) is significantly larger than that of TBPI (Engelmann et al., 1989; Olsson et al., 1989; Seckinger et al., 1989a). A hydropathy index computation (Kyte and Doolittle, 1982) of the deduced amino acid sequence of the protein (Figure IC) revealed two major hydrophobic regions (see roundended boxes in Figure 1D). One, at its NH2-terminus, is apparently the signal peptide whose most likely cleavage site





Residue No.

Fig. 1. Nucleotide sequence of the type I TNF receptor cDNA and the predicted amino acid sequence of the encoded protein. (A) The probes used or screening for the cDNA: (a) The NH₂-terminal amino acid sequence of TBPI. (b) Synthetic oligonucleotide probes, designed on the basis of the √H₂-terminal amino acid sequence, used for initial screening. (c) and (d) Probes overlapping with the probes presented in b, used to confirm the alidity of clones isolated in the initial screening. (B) Schematic presentation of the cDNA clores isolated from a human colon (C2) and from CEMimphocyte (E13) libraries and a diagram of the complete cDNA structure. Untranslated sequences are represented by a line. Coding regions are exed. The shaded portions represent the sequences which encode the signal peptide and the transmembrane domains. (C) Hydropathy profile of the redicted amino acid sequence of the TNF receptor. Hydrophobicity (above the line) and hydrophilicity (below the line) values were determined ing the sequence analysis software package of the University of Wisconsin genetic competer group (UWGCG) according to Kyte and Doolittle

Soluble and cell surface TNF receptors

_		
D.	1 CGGCCCAGTGATCTTGA	17
131	ACTODACATOTO ACTOTO TO TACTOTO ACTODACIONATICA ACTODACIONA ACTODACATA ACTODACATO ACTODAC	136
	-21 -10 -1 +1	255
	Not Gly Lou Har The Val Pro Asp Lou Lou Pro Lou Val Lou Lou Glu Lou Lou Vel Gly Ile Tyr Pro Ser Gly Val Ile Gly Lou	
256	ATG GGC CTC TCC ACC GTG CCT GAC CTG CTG CTG CTG CTG GTG CTC CTG GAG CTG TTG GTG GGA ATA TAC CCC TCA GGG GTT ATT GGA CTG	345
	10 120	,,,
	Val Pro His Leu Gly Asp Arg Glu Lys Arg Asp Ser Val Cys Pro Gin Gly Lys Tyr Ile His Pro Gln Asn Asn Ser Ile Cys Cys Thr	
346	GTC CCT CAC CTA GGG GAC AGG GAG AGA GAT AGE GTG TGT CCC CAA GGA AAA TAT ATC CAC CCT CAA AAT AAT TCG ATT TGC TGT ACC	435
	so	
	Lys Cys His Lys Oly The Tyr Lou Tyr Asn Asp Cys Pro Gly Pro Gly Gln Asp The Asp Cys Arg Clu Cys Glu Ser Gly Ser Pho The	
436	ANG THE CAE ANA EGA ACE THE THE TAE ANT GAE TOT CEA GGE CEG GGG CAE GAT ACE GAE TOE AGE GAE TOT GAG AGE GGE TEE TTE ACE	525
	70 90	
	Ala Ser Glu Ash His Lou Arg His Cys Lou Ser Cys Ser Lys Cys Arg Lys Glu Met Gly Gln Val Glu Ile Ser Ser Cys hr Val Asp	
526	GCT TOA GAA AAC CAC CTC AGA CAC TGC CTC AGC TGC TCC ANA TGC CGA AAG GAA ATG GGT CAG GTG GAG ATC TCT TCT TGC ACA GTG GAC	615 .
	_100 110	
	Arg Asp Thr Val Cys Gly Cys Arg Lys Asn Gln Tyr Arg His Tyr Trp Ser Glu Asn Leu Phe Gln Cys Phe Asn Cys Ser Leu Cys Leu	
616	CGG GAC ACC GTG TGT GGC TGC AGG AAG AAC CAG TAC CGG CAT TAT TGG AGT GAA AAC CTT TTC CAG TGC TTC AAT TGC AGC CTC TGC-CTC	705
	130	
	Asn Gly thr Vel His Leu Ser Cys Gln Glu Lys Gln Asn Thr Vel Cys Thr Cys His Ala Gly Phe Phe Leu Arg Glu Asn Glu Cys Val	
706	THE STORY OF	795
	Ser Cyal Ser App Cyal Live Live Ser Leu Civi Cyal The Live Ser Cyal Cyal Cyal Cyal Cyal Cyal Cyal Cyal	
796	The day for the day for the can he day was day for die kep Ser Gly for	
, 20	190	885
	The Val Lau Pro Lau Val Ile >he She Gly Lau Cys Lau Lau Ser Lau Lau Pro Lau Mat Tyr) Arg Tyr Gln Arg Trp Lys	
***	ACA GTG GTG TTG CCC CTG GTC ATT TTC TTT GGT CTT TGC CTT TTA TCC CTC CTC TTC ATT GGT TTA ATG TAT CGC TAC CAA CGG TGG AAG	
	220 230 24C	975
	Ser Lys Leu Tyr Ser Ile Val Cys Gly Lys Ser Thr Pro Glu Lys Glu Gly Glu Leu Glu Gly Thr Thr Thr Lys Pro Leu Ala Pro Asn	
976	TEC ANG CTC THE TEC ATT GTT TGT GGG ANA TEG ACA CET GAN ANA GAG GGG GAG CTT GAN AGA ACT ACT AND CEC CTG GEC CON AND	1065
	250 260 270	1003
	Pro Ser Phe Ser Pro Thr Pro Gly Phe Thr Pro Thr Leu Gly Phe Ser Pro Val Pro Ser Ser Thr Phe Thr Ser Ser Ser Thr Tyr Thr	
1066	CCA AGO TTO AGT COD ACT COA GGO TTO AGO COD AGO CTG GGO TTO AGT COD GTG CCC AGT TOO ACC TTO ACC TCC AGO TCC AGO TCA AGO	1155
	280 290 300	•
	Pro Cly Asp Cys Pro Asn Phe Ala Ala Pro Arg Arg Glu Val Ala Pro Pro Tyr Cln Gly Ala Asp Pro Ile Leu Ala Thr Ala Leu Ala	
1156	CCC GGT GAC TGT CCC AAC TTT CCG GCT CCC CGC AGA GAG GTG GCA CCA CCC TAT CAG GGG GCT GAC CCC ATC CTT GCG ACA GCC CTC GCC	1245
	320 330	
	Ser Asp Pro Ile Pro Asn Pro Leu Cln Lys Trp Glu Asp Ser Ala His Lys Pro Gln Ser Leu Asp Thr Asp Asp Pro Ala Thr Leu Tyr	
1246	THE GAD COD ATC COD AAC COD CIT CAG AAG THG GAG GAD AGG GCC CAC AAG CCA GAG AGG CTA GAD ANT GAT GAC COD GCG ACG CTG TAC	1335
	340 359 360	
	Ala Val Val Glu Asn Val Pro Pro Leu Arg Trp Lys Clu Pho Val Arg Arg Leu Gly Leu Ser Asp His Glu Ile Asp Arg Leu Glu Leu	
1336		1425
	370 380 290	
	Gin Ain Gly Arg Cys Lou Arg Glu Ala Gin Tyr Ser Het Lou Ala Thr Trp Arg Arg Thr Pro .= Arg Glu Ala Thr Lou Glu Lou	
1426		1515
	410 420	
1510	Leu Gly Arg Val Leu Arg Asp Het Asp Leu Leu 3ly Cys Leu Glu Asp Ile Glu Glu Ala Leu Cys Gly Pro Ala Ala Leu Pro Pro Ala	
1319		1605
	Pro Ser Jan Jan Ara Pad	
	Pro Ser Leu Leu Arg tod	
1719		1718
1838		1837
1957		1956
2076	Catargeagtitititig tettigtitigtitigtititiaa teaateagetgaretgarettiggaretgggaretgegaetggce <mark>eteggaeageagatageagatga</mark> Ctgteetaaggeggggggggggggaeggaacaatggggeetteagetggagetggagettitgtacataa <u>attaaaattetgaagttaaaaaaaaaa</u> 2176	2075
	2176	

(1982). The curve is the average of the hydrophobicity index for each residue over a window of nine residues. (D) Nucleotide and predicted amino acid sequences of the TNF receptor. The presumptive start and stop signals are denoted by asterisks; the three sequences derived from TBPI by broken overlining; the transmernbrane and leader domains by round-ended boxes; and the four repetitive sequences in the extracellular domain by thick underlining. Cysteine residues are boxed. Giycosylation sites are overlined and the presumptive polyadenylation signal is underlined.

Table I. Amino acid sequences of TBPI and TBPII

IBP i:																								_
CNBr-1 (=N-terminus) CNBr-2 C-terminus	NH;		p Se	r Va	l Cy l Gli	s Pro	o Gli	n Giv	8 Ly: r Cy:	s Tv	r 11 c	. ui	. D.		_	4 15 - ir Va								
•		100			1 00	OH		•																
IBP II:																					•			
N-terminus		1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	. 4	, ,					
TRP 35	-	. Ala	Gin	Val	Ala	Phe	Thr	Pro	Tyr	Ala	Pro	Glu	Pro	Giv	Se	The	C.,	, I.	, 10	3 19	9 20 9 Glu	21	22	
TRP 39/1	NH2	Leu	Cys	AJa	Pro	Leu	Arg	Lys									c,	. AI	g ce	U Ar	g Glu	Tyr	Tyr	•
TRP 39/2	NH ₂	Cys	Arg	Pro	Gly	Phe	Gly	Val	Ala	Arg														
TRP 44/1	NHS	Giu	Tyr	Tyr	Asp	Gin	Thr	Ala	Gin	Mot	Cue	Cys												
TRP 44/2	MLJ2	GiU	iyr	Tyr	Asp	Gir.	Thr	Ala	Gin	Met	C√s	Cvs	Ser											
	14115	301	CAR	G ₁ y	P/0	Ser	Tyr	Pro	Asp															
TRP 46/1	NH ₂	Phe	Thr	,oro	Tyr	Ala	Pro	Glu	Pro	GU	Sar	The	Cue	A										
TRP 46/2	NH ₂	Leu	Arg	Glu	Tyr	Tyr	Aso	Gin	The	Ala	Gin.	Mari	Cua	nry										
RP 50	NH ₂	Leu	Arg	Glu	Tyr	Tyr	Aso	Gir	The	Ala	Gin	Met	Cys	Cys	•••									
'RP 53/1	NH ₂	Val	Ala	Phe	Thr	Pro	Tur	Ala	Pro	Ciu	0:11	Mei	Cys	Cys										
RP 53/2	NH ₂	Cys	Arg	Pro	GIV	Phe	Gh/	Val	Ala	Ann	PIO	Giy	Se.	Thr	Cys	Arg						-		
RP 54/1	NH ₂	Pro	Glv	Tm	Tur	Cys	Ala	Lou	Car	Arg														
RP 54/2	NH ₂	Aia	Gin	Val	Ala	Pho	The	2	Jei Tur	Lys	_		_											
RP 60	NH ₂	lle	Cvs	The	Cre	Phe	Dea	Pro Oh	iyr T-	Ala	Pro	Giu	Pro	Gly	Ser	Thr	Cys	Arg						
RP 62	•		-,-		U 73	719	L(0	GIV	1 m	Tur	Cue	Ala	1	C										
AP 65	NH ₂ NH ₂	Pro	Gh	The	Ciu	TA:	Ser	ASP	Val	Val	Cys	Lys	Pro	Cys	Ala	Pro	Gly	Thr	Phe	Ser				
RP 67																								
RP 84																					l ve			
· •	NH ₂	1111	381	MSD	IN	٧a.	Cys	Asp	Ser	Cys	Giu	Asp	Ser	Thr	Tyr	Thr	Gin	Leu	Tro	-,3	-73			

(Von Heijne, 1986) lies between the glycine and isoleucine residues designated in Figure 1D as -1 and +1 respectively. The other major hydrophobic domain, located between residues 191 and 213, is flanked at both ends by several charged amino acids, characteristic of a membrane anchoring domain (Pidgeon et al., 1989). As in several other transmembrane proteins, the amino acids confining the hydrophobic domain at its COOH-terminal are basic. The transmembrane domain bisects the predicted protein into almost equally sized extracellular and intracellular domains.

The extracellular domain has three putative sites for asparagine-linked glycosylation (overlined in Figure 1D). Assuming that the amount of oligosaccharides in the extracellular domain is similar to that reported in TBPI (Seckinger et al., 1989b), the molecular size of the mature protein is very similar to that estimated for the type I receptor (~58 kd) (Hohmann et al., 1989; Engelmann et al., 1990a).

Features of the predicted amino acid sequence in the type I TNF-R and relationship to the structure of TBPI and TBPII

The amino acid sequence of the extracellular domain of the protein encoded by the E13 cDNA fully matches several determined TBPI amino acid sequences (Table I). It contains the NH₂-terminal amino acid sequence of TBPI at amino acids 20-32 (compare Figure 1D and 1A a), a sequence corresponding to the COOH-terminus of TBPI at amino acids 178-180, and also, adjacent to the first methionine located further downstream in the encoded protein, a sequence identical to the NH₂-terminal amino acid sequence of a cyanogen-bromide cleavage fragment of TBPI (broken lines n Figure 1D). There is also a marked similarity in amino cid composition between the extracellular domains of the eceptor and TBPI (Table II). The most salient feature of

Table II. Similarity of the amino acid compositions of the TNF binding protein TBPI and a corresponding region in the extracellular domain of the TNF-R (type I)

Amino acid	moi/100 mol	of amino acids
	TBPI°	Residues 20-180 in the extracellular domain ^b
Ala	1.7	1.2
Cys	12.8	14.9
Asp + Asn	10.9	11.1
Glu + Gln	13.9	12.4
Phe	3.2	3.1
Gly	6.3	5.6
His	4.4	4.3
lle	2.8	2.5
Lys	6.2	6.2
Leu	8.0	6.8
Met	0.4	0.6
Pro	3.8	3.1
Arg	4.7	4.3
Ser	8.1	9.3
Chr .	6.1	6.2
/al	4.2	4.3
ம்	-	0.6
`yr	2.4	3.1

[&]quot;According to Olsson et al., 1989.

this amino acid composition is a very high content of cysteine residues (shown boxed in Figure 1D). The positioning of the cysteine residues as well as of other amino acids within the extracellular domain displays a four-fold repetition pattern (Figure 2 and underlined in Figure 1D). As shown in Figure 2, there is a marked homology between this four-

Residue 20 corresponds to the NH₂-terminal amino acid of TBPI. Residue 180 is the COOH-terminal residue of TBPI.

Soluble and cell surface TNF receptors

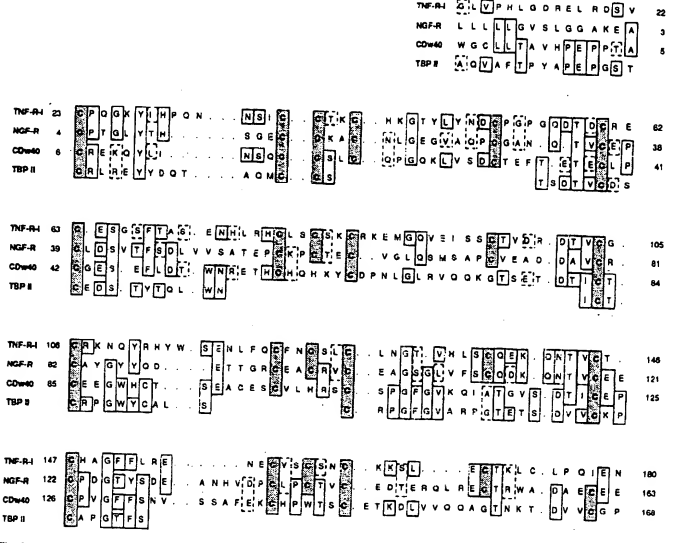


Fig. 2. Internal cysteine-rich repeats in the extracellular domain of the TNF-R and their alignment with the homologous repeats in the extracellular domains of human NGF-R and the CDw40 antigen and with sequences of amino acids present in TBPII. The amino acid sequences (one-letter symbols) are aligned for maximal homology. Dashes indicate gaps introduced to optimize the alignment. Identities in sequences are shown in boxes. Conservative substitutions (I = L = V; D = E; K = R = H; T = S; G = A; N = Q) are boxed with dotted lines. The positions of the amino acids within the receptors are denoted in the right and left hand margins. The sequences in TBPII (Table I and Materials and methods) were optimally aligned with the sequences of the other three protein; presented in this figure, using the 'Best Fit' alignment program in the UWGCG software package.

domain structure and sequences found in the extracellular domain of the receptor for the nerve growth factor (NGF-R) (Johnson et al., 1986; Radeke et al., 1987) and also, to a somewhat lower degree, to sequences in the extracellular domain of the recently cloned receptor-like B cell antigen CDw40 (Braesch-Andersen et al., 1989; Stamenkovic et al., 1989). The amino acid sequence within the extracellular domain of the TNF-R, which corresponds to the COOHterminal sequence of TBPI (see Table I and Figure 7), is located at the COOHterminus of the cysteine-rich repeat region. The sequence corresponding to the NH2-terminal sequence of TBPI corresponds to a sequence located a few amino acids upstream of the NH2-terminal end of this region (broken lines in Figure ID) in the extracellular domain.

In contrast to the identity of amino acid sequences between TBPI and the extracellular domain of the type I receptor, sequences examined in the soluble form of the type II TNF-R (TBPII, Table I) were not identical to any sequence in the type I TNF-R. This finding is expected, considering the

lack of immunological cross-reactivity between the two receptors (Engelmann et al., 1990b). However, as demonstrated in Figure 2, the sequences in TBPII have a significant homology of structure with the four-fold cysteinerich repeat region in the extracellular domains of the type I TNF-R, the NGF-R and the CDw40 protein. The similarity between TBPII and the CDw40 protein is particularly notable and seems to extend also to a region within the two proteins which protrudes from the cysteine-rich structures in the NH₂ direction (Figure 2).

In contrast to the very high content of cysteine residues in the putative extracellular domain of the type I TNF-R, there are only five cysteine residues in the intracellular domain. Between the two which are proximal to the transmembrane domain (positions 227 and 283) extends a stretch of 55 amino acids which is rich in proline residues (16% of the residues) and even richer in serine and threonine residues (36%), most located very close to or adjacent to each other. The consensus sequence Gly-x-Gly-x-x-Gly

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nucleotide binding proteins (Kamps et al., 1984) is not present in the intracellular domain.

Expression of the type I TNF-R cDNA

To explore the relation between the protein encoded by the E13 cDNA and TBPI further, we expressed this protein in CHO cells. The E13 cDNA was introduced into an expression vector and was cotransfected with a recombinant vector containing the dihydrofolate reductase (DHFR) cDNA into DHFR-deficient cells. After selection by growth in a nucleotide-free medium, individual clones were amplified by growth in the presence of methotrexate. A number of clones which react with several monoclonal antibodies that bind to spatially distinct epitopes in TBPI were detected (Figure 3). Expression of the protein was correlated with an increase in specific binding of human TNF to the cells (Table III).

Applying a sensitive immunoassay for TBPI, in which polyclonal antibodies and a monoclonal antibody against this protein were employed, we could also detect a soluble form of the protein in the growth medium of CHO cells, which

express the human TNF-R on their surface (Table III). All of five different CHO clones which expressed the TNF-R produced this soluble protein. Several other transfected clones which did not express the cell surface receptor did not produce its soluble form either (not shown). When analyzed by reversed phase HPLC, the CHO-produced soluble TNF-R eluted as a single peak, with a retention time identical to that of TBPI (Figure 4).

Northern blot analysis using the E13 cDNA as a probe

To gain information on the transcripts which encode the type I TNF-R, we tested mRNAs from cells of differing origin for their ability to hybridize with the E13 cDNA. As shown in Figure 5, in all the cell types, including the HT29 cells, which continuously secrete a soluble form of the type I TNF-R (Aderka,D., Nophar,Y., Engelmann,H. and Wallach,D., manuscript in preparation), only a single hybridizing transcript was detectable, in all cases of the same size i.e. -2300 bp, corresponding to the full length of the cDNA. Interestingly, significant amounts of this type

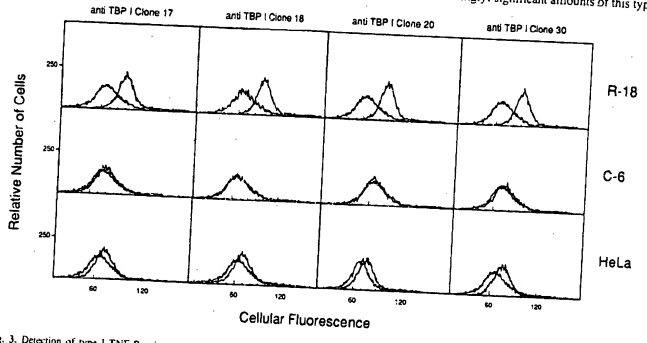


Fig. 3. Detection of type I TNF-R using monoclonal antibodies to TBPI in CHO cells transfected with E13 cDNA. CHO cells, clones R-18 (transfected with an expression vector in which the E13 cDNA was placed under the control of an SV40 promotor) and C-6 (control; a clone of cells transfected with an expression vector in which E13 was placed in the inverse orientation), and HeLa cells were stained with the anti-TBPI that observed when a mouse monoclonal antibody against TNF was used in the first step of the staining as a control.

CHO cell clone	he cell surface and soluble forms of human typ	THI THE CHO CEILS	
	Specific binding of TNF (c.p.m./10 ⁶ cells)	Cells expressing human cell surface TNF-R (% fluorescent cells)	Human soluble type TNF receptors (pg/ml)
Contransfected C6 -16 -18	180 ± 45 175 ± 50 550 ± 60 610 ± 40	<1% <1% 73% 89%	<0.03 <0.03 30

The R-16 and R-18 clones consist of cells transfected with a recombinant expression vector containing E13 cDNA. C-6 cells were transfected with a control vector (see Figure 3). Binding of radiolabeled TNF to the cells was determined in quintuplicate samples. Detection of immunoreactive receptors on the cells was carried out using combined 17, 18 and 20 anti-TBPI monoclonal antibodies. Results are expressed as percentage of fluorescent cells (background values, obtained by staining the cells with an anti-TNF monoclonal antibodies.

Soluble and cell surface TNF receptors

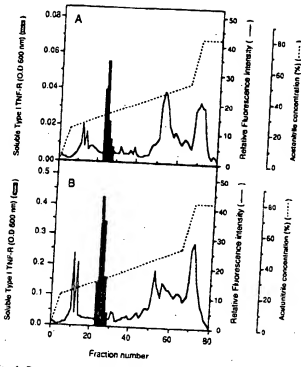


Fig. 4. Reversed phase HPLC of the CHO-produced, soluble form of the type I TNF-R. A concentrate of the conditioned medium of the CHO R-18 clones (see Figure 3) and a concentrate of the CHO C-6 clone to which 3 ng pure TBPI was added were applied to an Aquapore RP300 column. Elution was performed with a gradient of acetonitrile in 0.3% aqueous trifluoroacetic acid (---). Fractions were examined for content of protein (——) and of the soluble form of the type I by ELISA (EE), as described in Materials and methods. None of the eluted fractions of a concentrate of the CHO C-6 clone without addition of TBPI was found to contain any detectable amounts of the soluble form of the receptor (not shown).

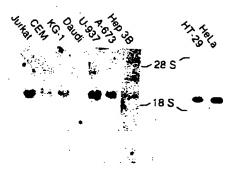


Fig. 5. Detection of the mRNA for the type I TNF-R by Northern blotting analysis. Hybridization of total RNA (25 µg/lane) from cells of the Jurkat (Gillis and Watson, 1980), CEM (Foley et al., 1965), KG-I (Koeffer and Golde, 1978), Daudi (Klein and Klein, 1968), K-562 (Lozzio and Lozzio, 1975), U-937 (Sundstroem and Nilsson, 1976), A-673 (Giard et al., 1973), Hep 3B (Aden et al., 1979), HT-29 (Fogh and Trempe, 1975), and HeLa (Gey et al., 1952) lines with the ³²P-labeled E13 insert was carried out as described in Materials and methods, 28S and 18S refer to ribosomal RNA size markers.

I TNF-R mRNA could also be detected in the U937 cells, in which the prevalent TNF-R is type II (Engelmann et al., 1990b), suggesting that post-transcriptional mechanisms take part in the control of the expression of the type I receptor.

Evidence for the involvement of type I TNF-R in stimulation of protein phosphorylation by TNF
Treating cells with TNF results in a rapid increase in the phosphorylation of certain specific cellular accession.

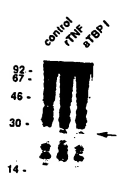


Fig. 6. Involvement of the type I TNF-R in stimulation of protein phosphorylation in cells. Effects of TNF- α (1000 U/ml) and rabbit antiserum to TBPI (1:1000) on the phosphorylation of proteins with a mol. wt of 27 kd in HeLa cells. Untreated cells served as a control. The 27 kd protein(s) are indicated with an arrow on the right and the migration of molecular weight markers (Amersham, UK) is shown on the left. Normal rabbit serum had no effect at a dilution of 1:1000 (not shown).

some with a mol. wt of ~27 kd (Hepburn et al., 1988; Kaur and Saklatvala, 1988; Schutze et al., 1989). Since it is apparent from the sequence data of the intracellular domain of the type I TNF-R that this receptor is devoid of intrinsic protein kinase activity, it was of interest to examine the extent to which this receptor is involved in TNF-mediated protein phosphorylation events. Antibodies to TBPI induce various effects in cells which are characteristic of TNF. This activity was shown to be correlated with the ability of the antibodies to cross-link the type I TNF-R molecules (Engelmann et al., 1990a). As shown in Figure 6, treating HeLa cells which express the type I TNF-R (Engelmann et al., 1990b) with antibodies to TBPI induced, as does TNF, a marked increase in the phosphorylation of protein(s) with a mol. wt of 27 kd, confirming that the type I TNF-R is involved in this effect.

Discussion

There is accumulating evidence for the natural occurrence of soluble forms of cell surface receptors. Such forms have been identified, for example, for the receptors to interleukin-2 (IL-2) (Rubin et al., 1985; Osawa et al., 1986). growth hormone (Leung et al., 1987), NGF (DiStefano and Johnson, 1988), interleukin-6 (Novick et al., 1989), interferon-y (Novick et al., 1989) and tumor necrosis factor (Engelmann et al., 1989, 1990b; Olsson et al., 1989; Seckinger et al., 1989a). Yet knowledge of the exact structure of these soluble receptors and of the mechanisms of their formation is still limited. The most thoroughly characterized so far is the soluble form of the 55 kd receptor for IL-2. Based on detailed sequence analysis and studies of its mode of formation in cultured cells, it was suggested that it is derived from the cell surface form of the receptor by proteolytic cleavage (Robb and Kutny, 1987).

A different mechanism for the formation of soluble receptors was proposed in two recent studies describing the cloning of the cDNAs for the receptors to IL-4 and IL-7. Besides cDNA clones encoding the full length receptors, clones which encode truncated, soluble forms of these receptors were also isolated in these studies. It was suggested that these latter clones are derived forms.

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transcribed from the same genes which encode the cell surface forms, but differently spliced (Mosley et al., 1989; Goodwin et al., 1990).

Data presented in our study are consistent with the notion that TBPI-the soluble form for the type I TNF-R-constitutes a fragment of the cell surface form of this receptor corresponding to its extracellular domain. The receptor is recognized by several monoclonal antibodies to TBPI which interact with several spatially distinct epitopes in this protein (the present study and Engelmann et al., 1990a). The amino acid sequence in the extracellular domain fully matches several sequences present in TBPI. Furthermore, the amino acid composition of the region within the extracellular domain which extends between those residues which correspond to the NH2- and COOH-termini of TBPI is very similar to the amino acid composition reported for TBPI. There is also a similarity in size between TBPI and this part of the receptor [taking into account that about a third of the TBPI molecule consists of oligosaccharides (Seckinger et ai... 1989b)]. Particularly informative with regard to the mechanism of formation of TBPI is the finding that a soluble form of the type I TNF-R is produced by CHO ceils transfected with the TNF-R cDNA. This finding implies that cells possess some mechanism(s) which allow(s) the formation of the soluble form of the TNF-R from the same transcript that encodes the cell surface form. There is no indication from the data of this study for the existence of transcripts which specifically encode soluble forms of TNF-R. Northern blot analysis did not reveal transcripts smaller than the full size of the TNF-R mRNA in any of the cells examined, not even in the HT29 cells, which continuously release significant amounts of a soluble form of the type I TNF-R into the culture medium. Furthermore, sequence and restriction mapping analyses of the various cDNA clones isolated in this study together with the C2 and E13 clones failed to reveal any difference in structure, besides differences in size, between these clones and the E13 cDNA (data not shown). The amino acid sequence data of TBPI also provide no indication of the existence of transcripts specific to this protein. Soluble receptors produced from alternatively spliced transcripts, as suggested for the II-4 and IL-7 receptors, are expected to have unique COOHterminal sequences (Mosley et al., 1989; Goodwin et al., 1990). The COOH-terminal sequence of TBPI was found to be identical to a sequence found in the cell surface receptor. Still, the existence of a minor population of transcripts which specifically code for soluble forms of TNF-Rs in amounts below the limit of detection of the techniques employed, although not supported by the data presented in this study, cannot be excluded.

The low rate of production of the soluble form of the type I TNF-R by the E13-transfected CHO cells does not necessarily reflect maximal activity. In HT29 cells, the spontaneous release of a soluble form of type I TNF-R occurs at about a 10-fold higher rate than that observed with the CHO-R-18 clone (data not shown). Furthermore, a recent study (Porteu and Nathan, 1990) indicates that the mechanism of formation of the soluble TNF-R can be effectively enhanced by certain specific stimuli. Stimulation of human neutrophils with N-formyl Met-Leu-Phe, or with several other physiological stimuli, was found to result, within a few minutes, in an extensive decrease of the cellsurface expressed TNF-R and an accompanying release of a soluble form of these receptors, similar in size to TBPI.

A likely mechanism whereby soluble forms of TNF receptors can be derived from the same transcripts which encode the cell surface forms is proteolytic cleavage. Indeed, flanking the amino acid residue which corresponds to the NH2-terminus of TBPI there are, within the amino acid sequence of the receptor, two basic amino acid residues (Lys-Arg) which can serve as a site of cleavage by trypsin-like proteases. The identity of the proteases which might cause cleavage to take place at the COOH-terminus of TBPI is not known. In view of the marked structural homology between the extracellular domain of the type I TNF-R and the soluble form of the type II TNF-R (TBPII) as well as the homology with the extracellular domain of the NGF-R, for which existence of a soluble form has been also documented (DiStefano and Johnson, 1988; Zupan et al., 1989), it is tempting to speculate that a common mechanism of cleavage and similar cleavage sites are involved in the formation of the soluble forms of those three receptors. Such a mechanism can have a dual effect on cell response to TNF. Its activation may result in the suppression of the response both in those cells in which it functions—as a consequence of the decrease in their intact cell surface receptors—as well as in other cells because of the ability of the released, soluble form of the receptor to sequestrate TNF. Detailed analysis of the biosynthesis of TNF-R, facilitated by use of cells transfected with TNF-R cDNA in vectors dictating its overexpression, should provide further information on the mechanism and functional implications of the formation of its soluble form.

Materials and methods

Determination of amino acid sequences within the TNF-binding proteins TBPI and TBPII

The TNF binding proteins TBPI and TBPII were isolated from concentrated preparations of urinary proteins, as described previously (Engelmann et al., 1990b) by ligand (TNF) affinity chromatography followed by reversed phase HPLC. TBPI was cleaved with cyanogen bromide, yielding two peptides which, following reduction and alkylation (Andrews and Dixon, 1987), were isolated by reversed phase HPLC. The two peptides (CNBr-1 and CNBr-2 in Table I) were subjected to NH2-terminal sequence analysis on a pulsed liquid gas phase protein microsequencer (Model 475A, Applied Biosystems Inc., Foster City, CA). The sequence found for one of the pepsides (CNBr-1) was identical to the NH2-terminal sequence of the intact TBPI protein (Engelmann et al., 1989, 1990b).

The COOH-terminal amine acid sequence of TBPI was determined by digestion of the protein with carboxypeptidase Y followed by sequential analysis of the released amino acids. A sample of pure TBPI (32 μ g) was mixed with I nmol norleucine, as an internal standard, dried thoroughly and resuspended in 8 µl 0.1 M sodium acetate buffer, pH 5.5, containing 0.8 μg carboxypepticlase Y (Sigma, St Louis, MO). Digestion was performed at room temperature. 2 μ l aliquots withdrawn at various time points were acidified by adding 3 μ l of 10% acetic acid to each, followed by addition of 15 μ l 0.5% EDTA. They were then subjected to automated amino acid analysis (Applied Biosystems, UK, mod. 420A). The results (Figure 7) indicate the sequence Ile-Glu-Asn-COOH.

Sequences within TBPII were determined by generation of tryptic peptides of the protein. A sample of pure TBPII (200 µg) was reduced, alkylated and repurified on an Aquapore RP-300 reversed phase HPLC column. Fractions containing the modified protein were pooled and the pH was adjusted to 8.0 with NaHCO3. Digestion with TPCK-trypsin (238 U/ing. Millipore Corp., Freehold, NJ) was performed for 16 h at room temperature at an enzyme to substrate ratio of 1:20 (w/w). The digest was loaded onto a C g RP-P reversed phase HPLC column (SynCrom, Linden, IN) and the poptides separated by a linear 0-40% acctonitrile gradient in 0.3% aqueous trifluoreacetic acid. The NH2-terminal amino acid sequences of the poptides and of the intact protein (N-terminus) are presented in Table I. The peptides were numbered according to their sequence of clutton from the RP-P column. In the fractions denoted as 39, 44, 46, 53 and 54, where heterogeneity of sequences was observed, both the major and the secondary sequences are presented.

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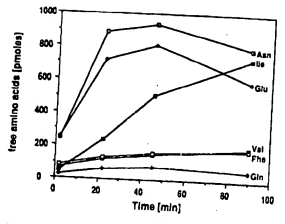


Fig. 7. Time course of the release of COOH-terminal amino acids from TBPI by carboxypeptidase Y.

Isolation of cDNA clones

Three mixtures of synthetic oligonucleotide probes generated from the nucleotide sequence deduced from the NH2-terminal amino acid sequence of TBPI were used for the screening of cDNA libraries. Initial screenings were carried out with 48-fold degenerate, 26-mers into which deoxyinosine was introduced, wherever the codon ambiguity allowed for all four nucleotides (Figure 1.A b). The validity of positive clones was examined by testing their hybridization to two mixed 17-mer nucleotide sequences, containing 96 and 128 degeneracies, corresponding to two overlapping amino acid sequences which constitute part of the sequences to which the 26 bp probes correspond (Figure 1A, c and d). An oligonucleotide probe corresponding to a sequence located close to the 5' terminus of the longest of the partial cDNA clones isolated with the degenerated proces (nucleotides 478-458 in Figure 1D) was applied for further screening cDNA libraries for a full length cDNA clone. ³²P-labeling of the probes, using T4 polynucleotide kinase, plating of the phages in lawns of bacteria, then screening them with the radiolabeled probes, isolation of the positive clones and subcloning of their cDNA inserts were carried out using standard procedures (Sambrook et al., 1989).

Nucleotide sequencing of the cDNA clones

cDNA inserts isolated from positive λ GT11 recombinant phages were subcloned into the pBluescript KS(-) vector. Inserts found in λ ZAP phages were rescued by excising the plasmid pBluescript SK(-) in them, using the R408 helper phage (Short et al., 1988). DNA sequencing in both directions was done by the dideoxy chain termination method (Sanger et al., 1977). Overlapping deletion clones of the cDNAs were generated, in both orientations, by digestion of the cDNA with exonuclease III ('Erase a base' kit, Promega Biotec, Madison, WI). Single-stranded templates derived from these clones using the R408 phage were sequenced with a T7 DNA polymerase sequencing system (Promega.)

Constitutive expression of the type I human TNF-R in CHO cells

The E13 insert was introduced into a modified version of the pSVL expression vector (kindly made available to us by Dr H.Kahana). This construct was transfected, together with the pSV2-DHFR plasmid which contains the DHFR cDNA, into DHFR deficient CHO cells (Chernajovsky et al., 1984), using the calcium phosphate precipitation method (Chen and Okayoma, 1987). Transfection with a recombinant pSVL vector which contained the E13 insert in the inverse orientation served as a control, Cells expressing the DHFR gene were selected by growth in nucleotide-free MEM α medium containing fetal calf serum which had been dialyzed against phosphate buffered saline. Individual clones were picked out and then further selected for amplification of the transfected cDNAs by growth in the presence of 500 nM sodium methotrexate.

Detection of surface-expressed type I TNF-R in the CHO cells Binding of radiolabeled human rTNF to cells (seeded in 15 mm tissue culture plates at a density of 2.5×10^5 cells/plate) was quantified as described before (Holtmann and Wallach, 1987).

To examine the binding of monocloral antibodies against TBPI to the CHO cells, the cells were detached by incubation in phosphate buffered saline (PBS: 140 mM NaCl, 1.5 mM KH₂PO₄, 8 mM Na₂KPO₄, 2.7 mM KCl, 0.5 mM MgCl₂, 0.9 mM CaCl₂), containing 5 mM EDTA and then incubated for 45 min at 0°C with 50 μg/ml of the test monocloral antibody

in PBS containing 0.5% bovine serum albumin, and 15 mM sodium axide (PBS/BSA). After washing the cells with PBS/BSA they were incubated further for 30 min at 0°C with FITC labeled, affinity purified goat antibody to the F(ab) fragment of mouse IgG (1:120 in PBS/BSA) (Bio-Makor, Israel) and then analyzed by determining the intensity of fluorescence in samples of 10⁴ cells using the Becton Dickinson fluorescence activated cell sorter 440. Three monoclonal antibodies to TBPI, clones 17, 18 and 20, shown by cross competition analysis to recognize four spatially distinct epitopes in the TBPI molecule (Engelmann et al., 1990a) and, as a control, a monoclonal antibody against TNF-or (all purified from ascitic fluids by ammonium sulfate precipitation and of the IgG₂ isotype) were used.

Quantification of the soluble form of the type I TNF-R by ELISA

A sensitive enzyme linked immunosorbent assay was set up, using TBPIspecific monoclonal and polyclonal antibodies in a sandwich technique. Inumunoglobulins of the anti-TBPI mAb clone 20 (Engelmann et al., 1990a) were adsorbed to 96 well ELISA plates (maxisorp, Nunc, Denmark) by incubation of the plates for 2 h at 37°C with a solution of 25 μ g/ml of the antibody in PBS. After incubating the wells further for 2 h at 37°C with n solution containing PBS, 1% BSA, 0.02% NaN3 and 0.05% Tween 20 (blocking solution) to block nonspecific further binding of protein, tested samples were applied in aliquots of 50 al/well. The plates were then incubated for 2 h at 37°C, russed 3 times with PBS supplemented with 0.05% Tween 20 (washing solution) and then rabbit polyclonal antiserum against TBPI, diluted 1:500 in blocking solution, was added to the wells. After further incubation for 12 h at 4°C, the plates were rinsed again and incubated for 2 h with horse radish peroxidase-conjugated purified goat anti rabbit IgG. The assay was developed using 2,2'-azino-bis (3-ethylbenzthiazoline-6 sulfonic acid) as a substrate (Sigma). The enzymatic product was determined colorimetrically at 600 nm. Pure TBPI served as a standard.

Detection of a soluble form of the type I TNF-R in the growth medium of the transfected CHO cells and its analysis by reversed phase HPLC

The amounts of the soluble form of the type I TNF-R in samples of the growth medium of the tested CHO cells, collected 48 h after medium replacement, were determined by the immunoassay described above. For analysis of the soluble receptor by reversed phase HPLC, the CHO cells were cultured for 48 h in serum-free medium (nucleotide-free MEM α). The medium samples were concentrated 100-fold by ultrafiltration on an Amicon PM10 membrane and 100 µl aliquots were then applied to an Aquapore RP300 column (4.6 × 30 mm, Brownlee Labs) pre-equilibrated with 0.3% aqueous trifluoroacetic acid. The column was washed with this solution at a flow rate of 0.5 ml/min until all unbound proteins were removed, and then eluted with a concentration gradient of acetonitrile in 0.3% aqueous trifluoroacetic acid, as described before (Engelmann et al., 1989). Fractions of 0.5 ml were collected and, after concentration in vacuo, were neutralized with 1 M HEPES buffer, pH 9.0. Amounts of soluble type I TNF-R in the fractions were determined by ELISA and the concentration of protein by the fluorescamine method (Stein and Moschera, 1981).

RNA isolation and analysis

RNA was isolated by a modification of the procedure described by Feramisco et al. (Feramisco et al., 1982; Queen and Baltimore, 1983) and analyzed by electrophoresis in 1.5% agarose/6% formaldehyde gels, followed by blotting to 'Genescreen plus' hybridization transfer membranes (NEN, Boston, MA). The E13 cDNA insert was ¹²P-labeled by random oligomer priming, using the Amersham random prinur labeling kit (Amersham, UK). The membranes were hybridized at 42°C in the presence of 50% formamide and then washed as prescribed by Sambrook et al. (1989) for the detection of low abundance sequences.

Determination of the effect of TNF and of antibodies to TBPI on protein phosphorylation

Confluent monolayers of HeLa cells (Gey et al., 1952), in 9 mm microwells, were incubated for 100 min with 100 μCi/ml [³²P]orthophosphate (Nuclear Research Center, Beer Sheva, Israel) in phosphate-free DMEM containing i6% fetal calf serum which had been dialyzed against 0.9% NaCl. Recombinant human TNF-or (kindly provided by Dr G. Adolf, Bochringer Institute, Vienna, Austria) and rabbit antiserum to TBPI (1:1000) (Engelmann et al., 1990b) or, for comparison, normal rabbit serum (1:1000) were then added to the cells for 20 min. The cells were rinsed, and immediately solubilized by boiling in SDS-PAGE sample buffer containing β-mercaptoethanol. The solubilized proteins were analyzed by SDS-PAGE

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Note added in proof

Since submission of this article, cloning of the cDNA for the type I TNF-R has been described in two publications [Loetscher, H., Pan, Y.-C.E., Lahm, H.W., Gentz, R., Brockhaus, M., Tabuchi, H. and Lesslauer, W. (1990) Cell, 61, 351-359 and Schall, T.J. et al., ibid., 361-370.] In two other publications [Smith, C.A. et al. (1990) Science, 248, 1019-1023 and Kohno, T. et al. (1990) Proc. Natl. Acad. Sci. USA (in press)] the cloning of the type II TNF-R was described The predicted amino acid sequence in the extracellular domain of this receptor fully matches the sequences presented here of amino acids in TBP II indicating further that, like TBPI, this soluble TNF binding protein is also derived from its immunologically cross-reacting cell surface TNF-R.

Appendix F

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Library	Vecto	mRNA or Source	# of Indep. Clones	Insert Size Range (Average)	Catalog Number
Human Libraries					
Human Adrenal cDNA	λgt10	Cortex and Medulla of adult with Cushing's Disease.	1.3 X 10 ⁶	0.55-3.3 kb (1.2 kb)	HL1014a
Human AML cDNA	λgt11	Acute myelogenous leukemia (KG-1).	1.4 X 10 ⁶	0.8–3.4 kb (1.3 kb)	HL1046b
Human B-Cell cDNA	Agt10	B-Cell leukemic cell line, RPMI 4265. Turner, et al., J. Biol. Chem., 250: 4512 (1975).	1.3 X 10 ⁶	0.39–3.4 kb (0.98 kb)	HL1018a
Human B-Cell cDNA	λgt11	B-Cell leukemic cell line, RPMI 4265. Turner, et al., J. Biol. Chem., 250: 4512 (1975).	8.9 X 10 ⁵	0.73-4.1 kb (1.2 kb)	HL1018b
Human B-Cell, PMA activated, cDNA	λgt10	RPMI 4265 cells treated with 50 ng/ml Phorbolmyristate acetate PMA for 4 days.	1.4 X 10 ⁶	0.6-3.7 kb (1.2 kb)	HL1035a
Human Brain cDNA	λgt11	Normal temporal cortex tissue, excised from around a tumor.	7.3 X 10 ⁶	0.4–3.1 kb (780 bp)	HL10036
Human Brain, Alzhelmer's, oONA	Agt11	Tissue around the hippocampus of a 70-year old diseased female, removed 5-8 hours after death.	8.1 X 10 ⁵	0.31-3.1 kb (1.1 kb)	HL1028b
Human CML Spleen cDNA	λgt11	Spleen of eight year old female with Chronic Myelogenous Leukemla.	1.4 X 10 ⁶	0.5–3.8 kb (1.1 kb)	HL1040b
Human Colon cDNA	λgt10	Normal tissues excised from around the colon cancer of a 53 year old male.	1.5 X 10 ⁶	0.6-3.2 kb (0.9 kb)	HL1034a
Human Colon cDNA	λgt11	Normal tissues excised from around the colon cancer of a 53 year old male.	1.3 X 10 ⁶	0.6-3.5 kb (1.1 kb)	HL1034b
Human Endothelial cONA	λgt11	Endothelial cells from umbilical cord velns. Cells were serially passaged and poly(A) isolated.	1.5 X 10 ⁶	0.34–3.4 kb (0.9 kb)	HL1024b
Human Epitheliai cDNA		Normal thymus epithellal cells, excised during open-heart surgery on a 3 year old Caucasian female.	1.4 X 10 ⁶	0.42-2.8 kb (0.9 kb)	HL1025b

Library	Vect	mRNA or Source	# of indep. Clones	Range	Cataion
Human Erythroleukemik cDNA	kgt10	Human erythroleukemic K562 cells cultured in RPM 1640. Slebert and Fukuda, J. Biol. Chem., 260: 640 (1985). Lozzlo and Lozzlo, Blood. 45: 321 (1975).	•	0.5-3.4 kb (1.0 kb)	HL1032a
Human Eye cDNA	λ gt10	Male/lemale eye pool.	1.1 X 10 ⁵	0.4–3.2 kb (0.7 kb)	HL1047a
Human Eye cDNA	λ gt11	Male/female eye pool.	1.5 X 10 ⁶	0.4-2.8 kb (0.7 kb)	HL1047b
Human Fibroblast, Lung, cDNA	λgt11	Lung fibroblast cell line, IMR-90. ATCC# CCL185.	2.2 X 10 ⁶	0.5-3.0 kb (1.3 kb)	HL1011b
Human Fibroblast, Skin, cDNA	λ gt10	Cultured primary fibroblasts from a young male.	1.4 X 10 ⁸	0.5-3.6 kb (1.0 kb)	HL1052a
Human Fibroblast, Skin, cDNA	λ gt11	Cultured primary fibroblasts from a young male.	1.1 X 10 ⁶	0.5-3.5 kb (1.1 kb)	HL1052b
Human Fibrosarcoma cDNA	λ gt11	HT-1080 tumor oells. Rasheed et al., Cancer 33:1027 (1974). ATCC # CCL121.	1.6 X 10 ⁸	1.0–3.4 kb (1.3 kb)	HL10485
Human Glioma cDNA	λ gt11	HS 683 from explant cultures of a glioma taken from the left temporal lobe. ATCC # HTB 138.	1.1 X 10 ⁶	0.4–3.1 kb (0.9 kb)	HL1049b
łuman Hairy Ceil aukemia (Mo-B) cDNA	λgt10		1.4 X 10 ⁶	0.4–3.2 kb (1.0 kb)	HL1043a
luman Heart cDNA		Adult male heart, including some aorta region. Poly(A) RNA was slightly degraded as visualized on alkaline agarose gel.	1.27 X 10 ⁵	0.4-3.4 kb (0.9 kb)	HL1038b
uman Heart, Fetal orta, cONA		Normal, third trimester heart	8.4 X 10 ⁵	0.5-3.5 kb (1.1 kb)	HL1042b
uman Hela Cell cona	1 3 1.	HeLa-derived D98-AH2 calls, HPRT- in phenotype. W.S. Szybalski et al., Natl. Cancer Institute Monograph, 7: 75 (1962).	1.3 X 10 ⁸	0.48-3.1 kb (0.86 kb)	HL1022b
man Hepatoma cDNA	Agt11 F	depatoma cells, G2.	1.3 X 10 ⁸	0.5-3.7 kb (0.96 kb)	HL1015b

Library	Vec	mRNA tor Source	# of Indep. Clones	insert Size Range (Average)	Catalon
Human Keratinocyte cDNA	λgt11	Primary keratinocyte cultur from adult epidermis.	re 1.7 X 10 ⁶	0.5-3.8 kb (1.1 kb)	HL1045b
Human Kidney oDNA	Agt10	From a Juvenile male whos kidney had been perfused for 24 hours prior to poly(A RNA isolation.		0.5-3.2 kb (0.9 kb)	HL1033a
Human Kidney cDNA	λgt11	From a juvenile male whose kidney had been perfused for 24 hours prior to poly(A) RNA isolation.		0.4-3.1 kb (0.9 kb)	HL1033b
Human Leukocyte Genomic	EMBL	3 Leukocyte genomic DNA, male.	8.8 X 10 ⁵	16kb	HL1006d
Human Leukocyte, Peripheral Blood, cDNA	λgt10	Peripheral blood leukocytes from adult female with Acute Promyelocytic Leukemia, HL60. S.J.Collins et al., Nature, 270: 347 (1977). ATCC# CCL240.	7.2 X 10 ⁵	0.69-3.5 kb (0.88 kb)	HL1020a
Human Leukocyte, Peripheral Blood, cDNA	λgt11	Peripheral blood leukocytes from adult female with Acute Promyelocytic Leukemla, HL60. S.J.Collins et al., Nature, 270: 347 (1977). ATCC# CCL240.	8.4 X 10 ⁵	0.77–3.6 kb (1.1 kb)	HL1020b
Human Liver cDNA	λ g t10	Normal adult liver, female.	1.5 X 10 ⁵	0.17-2.4 kb (0.8 kb)	HL1001a
Human Liver cDNA	λgt11	Normal adult liver, female.	7.8 X 10 ⁵	0.3-2.9 kb (0.94 kb)	HL1001b
Human Liver, Fetal, cDNA	λgt11	Fetal liver, 1st trimester, male.	2.3 X 10 ⁵	0.14–2.3 kb (1.1 kb)	HL1005b
Human Lung cDNA	λgt11	Normal adult lung tissue, excised during surgery.	8.2 X 10 ⁵	0.25-3.1 kb (1.2 kb)	HL1004b
luman Lung Fibroblast DNA	λgt11	Lung fibroblast cell line, IMR-90. ATCC# CCL186.	2.2 X 10 ⁶	0.5-3.0 kb (1.3 kb)	HL1011b
luman Lung Small Cell arcinoma (NCI-H69) DNA	λgt10	Lung Small Cell Carcinoma cell culture, NCI-H69.	1.4 X 10 ⁵	0.6-3.6 kb (0.93 kb)	HL1012a
uman Lung Small Cell arcinoma (NCI-H69) DNA	λgt11	Lung Small Cell Carcinoma cell culture, NCI-H69.	7.7 X 10 ⁵	0.4-2.8 kb (1.0 kb)	HL1012b

Library	Ved	mRNA stor Source	# of indep. Clones	insert Size Range (Average)	Catalog
Human Lung Small Cel Carcinoma (NCI-H128) cDNA		From NCI-H128 cell line derived from malignant pleural fluid of a 60 year or Black male. Gazder, et al. Cancer Research, 40: 350 (1980).	1	0.3-3.5 kb (0.85 kb)	HL10308
Human Lung Small Cell Carcinoma (NCI-H128) cDNA	Agt11	From NCI-H128 cell line derived from mallgnam pleural fluid of a 50 year ole Black male. Gazder, et al., Cancer Research, 40: 3502 (1980).		0.3–3.7 kb (0.92 kb)	HL1030b
Human Lung, WI-38, cDNA	λgt11	WI-38 human diploid cell line, derived from normal Caucasian female lung. L Hayflick, Exp. Cell Res., 25: 585 (1961).	1.2 X 10 ⁸	0.6–3.5 kb (1.1 kb)	HL1041b
Human Lymphocyte cDNA	Agt10	Near confluent Raji cells, B lymphocytes. Pulvertaft, Lancet, 1: 238 (1964). ATCC# CCL86.	1.5 X 10 ⁶	0.25-3.2 kb (0.87 kb)	HL1002a
Human Mammary Gland cDNA	λ g t10	Adult female breast tissue excised during masteotomy. The female was in the 8th month of pregnancy, showing well-differentiated tissues and lactational competence.	1.1 X 10 ⁸	0.6-3.3 kb (1.1 kb)	HL1037a
Human Mammary Gland CDNA	λgt11	See above.	1.1 X 10 ⁶	0.5–3.4 kb (1.2 kb)	HL1037b
Human Melanoma cDNA	λgt11	Near confluent melanoma A2058 cells.	7.1 X 10 ⁵	0.3–3.6 kb (0.82 kb)	HL1023b
Human Monocyte, Peripheral Blood, cDNA	λ gt 10	90% human peripheral blood monocytes, LPS-activated.	1.34 X 10 ⁶	0.4-3.5 kb (1.2 kb)	HL1050a
duman Monocyte, Peripheral Blood, cDNA	λ gt11	90% human peripheral blood monocytes, LPS-activated.	1.3 X 10 ⁶	0.5–3.7 kb (1.0 kb)	HL1050b
luman Monocyte, HP-1, cDNA		THP-1 monocytes from 1 year old male with Acute Monocytic leukemia. Int. J. Cancer, 26: 171 (1980). Cancer Research, 42: 1530 (1982). ATCC# TIB202.	1.4 X 10 ⁶	0.65–3.0 kb (0.97 kb)	HL1021a

Library	Vector	mRNA Source	# of Indep. Clones	Insert Size Range (Average)	Catalog Number
Human Monocyte, U-937, cDNA	λgt10	U-937 cultured cells actively growing prior to poly(A) RNA isolation.	1.5 X 10 ⁶	0.4-3.4 kb (0.95 kb)	HL1029a
Human Monocyte, U-937, cDNA	λgt11	U-937 cultured cells actively growing prior to poly(A) RNA isolation.	1.2 X 10 ⁶	0.3–3.7 kb (0.95 kb)	HL1029b
Human Monocyte, U-937, PMA activated cDNA	λgt10	U-937 cells treated with 50 ng/ml Phorbolmyristate acetate (PMA) for 3.5 days to achieve monocyte-like stage.	1.4 X 10 ⁵	0.6-3.8 kb (1.1 kb)	HL1036a
Human Monocyte. U-937, PMA activated cDNA	λ g t11	U-937 cells treated with 50 ng/mi Phorbolmyristate acetae (PMA) for 3.5 days to achieve monocyte-like stage.		1.0–3.9 kb (1.3 kb)	HL1036b
Human Multiple Myeloma cDNA	λgt11	Bone marrow obtained from a female with Multiple Myeloma, IM9. Ann. N.Y. Acad. Scl., 190: 221 (1972). PNAS, 71: 84 (1974). J. Biol. Chem., 249: 1661 (1974). ATCC# CCL159.	2.2 X 10 ⁶	0.52–3.5 kb (1.1 kb)	HL1027b
Human Neuroblastoma cDNA	λgt10	Neuroblastoma cell line, Kelly.	1.05 X 10 ⁵	0.4-3.4 kb (1.3 kb)	HL1007a
Human Osteosarcoma cDNA	Agt11	Osteosarcoma cell culture, MG-63. Antimicrob. Ag. Chemother., 12: 11 (1977). ATCC# CRL1427	1.6 X 10 ⁶	0.55–3.0 kb (0.97 kb)	HL1013b
Human Pancreas cDNA	Agt10	Islets of Langerhans	1.0 X 10 ⁶	1.2–4.0 kb (2.0 kb)	HL1054a
Human Peripheral Blood Leukocyte cDNA	Agt10	Peripheral blood leukocytes from adult female with Acute Promyelocytic Leukemia, HL60. S.J.Collins et al., Nature, 270: 347 (1977). ATCC# CCL240.	7.2 X 10 ⁵	0.69-3.5 kb (0.88 kb)	HL1020a
Human Peripheral Blood Leukocyte cDNA	λgτ11	Peripheral blood leukocytes from adult female with Acute Promyelocytic Leukemia, HL60. S.J.Collins et al., Nature, 270: 347 (1977). ATCC# CCL240.	8.4 X 10 ⁵	0.77–3.6 kb (1.1 kb)	HL1020b
Human Peripheral Blood Monocyte cDNA		90% human peripheral blood monocytes, LPS-activated.	1.34 X 10 ⁶	0.4–3.5 kb (1.2 kb)	HL1050a

Library	Ve	ector	mRNA Source	# (Indep Clone	D. Bano	8 Catala
Human Peripheral Blo Monocyte cDNA	od λg	ηt 11	90% human peripheral blood monocytes, LPS-activated.	1.3 X 1		sb HL1050b
Human Peripheral Bloc Monocyte cDNA	3		90% human peripheral blood monocytes.	1.2 X 10	0.0 · 0.4 A	
Human Placenta cona	λgti	13	Placental tissue, 34 week old.	1.0 X 10	U.U U.U KI	
Human Prostate cDNA	λgt		Normal 65 year old prosta	ate 1.5 X 10 ⁶	70	HL1051a
Human Prostate cDNA	λ gt1		formal 65 year old prosta	te 1.62 X 10 ⁶	U.S G.O KD	HL1051b
Human Retina cDNA	λgt1() A	dult retina.	1.6 X 10 ⁶	0.0 AD	HL1055a
Human Retina cDNA	λgt11	Ac	duit retina.	1.4 X 10 ⁸	(1.1 kb) 0.5–3.4 kb	HL1055b
Human Skin Fibroblast cDNA	λ gt10		iltured primary fibroblasts m a young male.	1.4 X 10 ⁵	(1.0 kb) 0.5–3.6 kb	HL1052a
Human Skin Fibroblast cDNA	λ gt11	_	itured primary fibroblasts m a young male.	1.1 X 10 ⁶	(1.0 kb) 0.5–3.5 kb	HL1052b
Human Spleen cDNA	λgt10	No	rmai 25 year old male een.	1.4 X 10 ⁶	(1.1 kb) 0.6–3.4 kb	HL1039a
Human Spieen cDNA	λgt11	Nor sple	mal 25 year old male en.	1.3 X 10 ⁵	(1.0 kb) 0.6–3.3 kb	HL1039b
Human Spleen, CML, cDNA	λgt11	WILL	en of 8 year old female Chronic Myelogenous remia.	1.4 X 10 ⁶	(1.0 kb) 0.5–3.8 kb (1.1 kb)	HL1040b
Human Submaxillary Gland cDNA	λ gt11	Male blops	gland excised during	1.3 X 10 ⁶	0.5-3.2 kb	HL1053b
Human T-Cell cDNA	Agt10	Jurka J. Car Yanaç	l leukemic cell line. it, Schnelder et al., Int. ncer, 19:621 (1977). gi et al., PNAS, 82: (1985).	1.1 X 10 ⁸	(1.0 kb) 0.58–3.8 kb (1.2 kb)	HL1016a
Human T-Cell cDNA	λ g t11	T-Cell Jurkat J. Can	Laukemic cell line. Schnelder et al., Int. cer, 19:621 (1977). I et al., PNAS 82	1.1 X 10 ⁶	0.58-3.8 kb (1.2 kb)	HL1016b

Library	V	ector	mRNA Source		# d Inder Clone	o. ¨	nsert S Rang Averag	•	Catalog Number
Human T-Cell, PHA Stimulated, cDNA	λς	P	Concentrated T-Cell topulation from periphe lood of a healthy adult. HA stimulated for 48 ho		1.5 X 10) ⁶ 0	.45-3.8) (0.95 kb)	 ф	HL1031a
Human T-Cell, PHA Stimulated, cDNA	λg:	111 C po bl	oncentrated T-Cell pulation from periphen ood of a healthy adult. A stimulated for 48 ho	ai	1.0 X 10	U	.7-3.3 kt (1.4 kb))	HL1031b
Human Testis cDNA	λgti	11 No du hei	ormal testicle, excised ring surgery from a althy 50 year old. Not monally induced.		1.0 X 10 ⁶	Ψ.	7–3.3 kb 1.2 kb)		HL10105
Human Thymocyte DNA	λgt1	1 Noi dun	mal thymus excised ing surgery on a 3 year Caucasian female.		1.7 X 10 ⁶		5–3.4 kb l.9 kb)		dL1026b
Human Thyroid Carcinoma cDNA Human Wilms' Tumor	λgt11	Adu	it thyroid carcinoma	6	3.8 X 10 ⁵	0.65 /1	-2.7 kb 2 kb)	н	L1009b
(G-401) cDNA	λgtī 1			1	.3 X 10 ⁵		3.8 kb	<u>t</u>	L1044b
Monkey Librarian							J.U KU	TI.	-10440
Monkey Libraries Monkey Brain cDNA	λgt10	Adult	Male Phonus hast.					п	-10440
Monkey Brain cDNA	λgt10	Adult tissue	male Rhesus brain	1.3	3 X 10 ⁶	0.5-3			1003a
Monkey Brain cDNA Monkey Brain cDNA	λ g t10 λgt11	แระบอ	male Rhesus hrain		3 X 10 ⁶ 2 X 10 ⁵		.4 KD	OL	1003а
Monkey Brain cDNA Monkey Brain cDNA Monkey Genomic		Adult i tissue.	male Rhesus hrain	1.2		0. 5 –3 0.6–3.	.4 kb 7 kb kb	OL OL1	
Monkey Brain cDNA Monkey Brain cDNA Monkey Genomic	λgt11	Adult tissue. Adult f	male Rhesus brain	1.2	! X 10 ⁵	0.5-3 0.6-3 10-22 (16.3	.4 kb 7 kb kb kb)	OL OL1	1003a 1003b 004d
Monkey Brain cDNA Monkey Brain cDNA Monkey Genomic Monkey Liver cDNA Monkey Liver cDNA	λgt11 EMBL3	Adult it tissue. Adult if Adult in tissue.	male Rhesus brain emale Rhesus liver.	1.2 1.5	X 10 ⁵ X 10 ⁶ X 10 ⁵	0. 5 –3 0.6–3.	.4 kb 7 kb kb kb)	OL1 OL1	1003a 1003b 004d
Monkey Brain cDNA Monkey Brain cDNA Monkey Genomic Monkey Liver cDNA Monkey Liver cDNA	Agt11 EMBL3 Agt10 Agt11	Adult in tissue. Adult in tissue. Adult in tissue. Adult in tissue. Adult in tissue.	male Rhesus brain emale Rhesus liver nale Rhesus liver ale Rhesus liver	1.2 1.5	X 10 ⁵ X 10 ⁶ X 10 ⁵ X 10 ⁶	0.5-3 0.6-3 10-22 (16.3 0.5-3.4	.4 kb 7 kb kb) kb	OL OL1	1003a 1003b 004d 002a

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Protocols and Maps

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CLONTECH Ordering Information

Technical Information:

Our scientific staff is happy to assist you with any questions concerning the use of our products.

CLONTECH is committed to providing you with the best innovative products and services possible. We value you as a customer. If you are dissutisfied in any way with our products or services, please let us know.

We are easy to reach between the hours of S AM and 6 PM, Monday through Friday at:

(415) 424-8222

or, you may contact us through your distributor.

How to order:

Outside the United States:

CLONTECH's distributors around the world are pleased to accept your orders for their countries of license. Please refer to the distributor list in this appendix for contact information. If you are profering directly from Canada, please supply the name and telephone number of your Customs broker.

Within the United States:

To expedite your order, please provide the following information:

- 1. Shipping address
- 2. Billing address
- 3. Purchase order number
- 4. Catalog number and product description
- 5. Size and quantity of product
- 6. Purchaser's name and contact number

Orders are processed upon receipt, in most cases within 24 hours for routinely inventoried products.

To avoid duplication of your order, please make sure that written confirming orders are clearly marked as such. CLONTECH does not require written confirmation of phone orders. If you receive duplicate shipments, please contact us immediately

Telephone orders:

Call (800) 662-CLON (outside CA) or (415) 424-8222 (inside CA)

Fax orders:

Our fax number is: (415) 424-1352

Telex orders:

Our telex number is: 330060

Mail orders:

Please send your ordering information to:

CLONTECH Laboratories, Inc.
4030 Fablan Way
Palo Alto, CA 94303
Attention: Customer Service Department

Standing Orders:

CLONTECH offers periodic shipments on any purchase order as a service to our frequent customers. Please arrange a schedule with your customer service representative.

Custom and Bulk Orders:

Please request quotations for all custom and bulk orders from your Customer Service Representative.

Minimum Order:

CLONTECH has no minimum order quantity. There is a \$5.00 handling charge for orders under \$150.00.

CLONTECH Ordering Information

Pricing and Terms:

Prices are subject to change without netice. For the most up-to-date pricing information on items in this price listing, please contact us or your distributor.

In the event of interim price changes on products amounting to more than 10% on orders over \$100.00, CLONTECH will contact you prior to shipping your order. Payment terms are not 30 days in US dollars.

US Government Customers

CLONTECH is listed on the GSA Schedule. Our NIH BPA Number is 263-00025536.

Shipping:

CLONTECH products are shipped F.C.3. Palo Alto, CA. Shipping charges are prepaid and added to your invoice. Orders requiring blue or dry ice are shipped via UPS Next Day Air or Federal Express P1. Monday through Thursday, unless otherwise requested. Orders not requiring ice are shipped via UPS 2 Day Air, Monday through Friday. Shipments of hazardous items are sent by an appropriate carrier.

Alternative carriers may be requested with your order.

Partial Shipments

Normally, all orders may be filled within a single shipment. If this is not possible, we will initiate a partial shipment of your order. If you do not want a partial shipment, please indicate this when you order.

Returns and Credit:

In order to facilitate processing, we request that no returns be made without prior authorization. Please contact your customer service representative for instructions for all returned goods. In order to receive credit, products should be returned in original, intact condition. There will be a 20% restocking fee on all goods returned due to customer ordering error.

Conditions:

All of CLONTECH's products are intended to be used only in a research laboratory. They are not to be used either for drug or diagnostic purposes, nor are they intended to be for human use.

Product Changes:

We reserve the right to delete products or change specifications at any time without notice.

Genetic Research Guidelines:

We recommend that our products be used in accordance with NIH guidelines developed for genetic research.

Patent Information:

No license or immunity under any patent is either granted or implied by the sale of any of our products.

Disclaimer:

CLONTECH disclaims any and all responsibility for any injury or damage which may be caused by the failure of the buyer or any other person to use these products in accordance with the conditions outlined herein.

Warranty:

CLONTECH's products are warranted to meet our product specifications in effect at the time of shipment. Notice of non-conforming products must be made to CLONTECH within 30 days of receipt of the product. This warranty is exclusive and is in lieu of all other warranties, expressed or implied, including any implied warranty of merchantability or fitness for any particular purpose. CLONTECH shall not be liable for any incidental, consequential or contingent damages.

Pricing and Terms:

Prices are subject to change without notice. For the most up-to-date pricing information on items in this price listing, please contact us or your distributor.

In the event of interlm price changes on products amounting to more than 10% on orders over \$100.00, CLONTECH will contact you prior to shipping your order. Payment terms are net 30 days in US dollars.

US Government Customers

CLONTECH is listed on the GSA Schedule. Our VA contract number is V797P-5824K. Our NIH BPA Number is 263-00025536.

Shipping:

CLONTECH products are shipped F.O.B. Palo Alto. CA. Shipping charges are prepaid and added to your invoice. Orders requiring wet or dry ice are shipped via UPS Next Day Alt or Federal Express P1, Monday through Thursday, unless otherwise requested. Orders not requiring ice are shipped via UPS 2 Day Air, Monday through Friday. Shipments of hazardous items are sent by an appropriate carrier.

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We recommend that our products be used in accordance with NIH guidelines developed for genetic research.

Patent Information:

No license or immunity under any patent is either granted or implied by the sale of any of our products.

Disclaimer:

CLONTECH disclaims any and all responsibility for any injury or damage which may be caused by the failure of the buyer or any other person to use these products in accordance with the conditions outlined herein.

Warranty:

CLONTECH's products are warranted to meet our product specifications in effect at the time of shipment. Notice of non-conforming products must be made to CLONTECH within 30 days of receipt of the product. This warranty is exclusive and is in lieu of all other warranties, expressed or implied, including any implied warranty of merchantability or fitness for any particular purpose. CLONTECH shall not be liable for any incidental, consequential or contingent damages.

Library	Vecto	mRNA ir Source	# of Indep. Clones	Insert Size Range (Average)	Catalon
Hamster Librarie	es				
CHO-K1 CDNA. 5'-STRETCH	λgt10	Chinese hamster ovary CHO-K1 cells. ATCC#CCL61.	1.4 X 10 ⁶	0.6 to>4.0 kb (1.5 kb)	JL1001a
A. Sissing CH	Àgt11	Chinese hamster ovary CHO-K1 cells. ATCC#CCL61.	1.5 X 10 ⁶	0.7 to>4.0 kb (1.5 kb)	JL10015
CHO-K1 cDNA, 5'-STRETCH	SWAJ-2	2 See above.	1.4 X 10 ⁶	0.6 to>4.0 kb (1.1 kb)	JL100th
Human Libraries		•			
al cDNA	λgt10	Cortex and Medulla of adult with Cushing's Disease.	1.3 X 10 ⁶	0.55-3.3kb (1.2 kb)	HL1014a
ruman AML cONA	λgt11	Acute myelogenous leukemia(KG-1).	1.4 X 10 ⁶	0.8–3.4 kb (1.3 kb)	HL1046b
Heman, Aorta, Fetal, IONA	λ g :11	Normal, third trimester, male aortal tissue.	8.4 X 10 ⁵	0.5–3.5 kb (1.1 kb)	HL10425
`-TcDNA	λ g t10	B-Cell leukemic cell line. RPMI 4265. Turner, et al., J. Biol. Chem., 250: 4512 (1975).	1.3 X 10 ⁶	0.39–3.4kb (0.98 kb)	HL1018a
łuman B-Cell cDNA	lgt11	See above.	8.9 X 10 ⁵	0.73-4.1kb (1.2 kb)	HL1C18b
ruman B-Cell, 1965-1777 /ated, cDNA		RPMI 4265 cells treated with 50 ng/ml Phorbolmyristate acetate PMA for 4 days.	1.4 X 10 ⁵	0.6-3.7 kp (1.2 kb)	HL1035a
uman Bone Marrow, -STRETCH cDNA		Adult male.	1.4 X 10 ⁶	0.6 to>4.0 kb (1.5 kb)	HL1058a
-STRETCH cDNA	λgt11 /	Adult male.	1.51 X 10 ⁶	0.6 to>4.0 kb (1.6 kb)	HL1058b
eman Brain, Tydda:acDNA	λ g t10		1.6 X 10 ⁶	0.6-3.5 kb (1.2 kb)	HL1080a
mygdala cDNA	λgt11		1.3 X 10 ⁶	0.6-3.7kb (1.2kb)	HL1080b
man Brain, Calcarine	λ g τ10		1.4 X 10 ⁶	_	HL:081a
man Brain, Calcarine Cex cONA	λgt11	562-CLON (outside CA)	1.0 X 10 ⁸		HL108:5

Library	Vector	mRNA Source	# of Indep. Clones	Insert Size Range (Average)	Catalog Number
Human Brain, Caudate cDNA	λgt10		1.8 X 10 ⁶	0.6-3.8 kb (1.4 kb)	HL1076a
Human Brain, Caudate cONA	λgt11		1.5 X 10 ⁶	0 6-3.8kb +1 4 kb)	HL1076b
Human Brain, Cerebellar Vermis cDNA	10ود		1.8 X 10 ⁶	0 6-3.7kb (1.3 kc)	HL1083a
Human Brain, Cerebellar Vermis cDNA	λgt11		1.2 X 10 ⁶	0 6–3.6 kb (1.2 kb)	HL1083b
Human Brain, Cingulate Gyrus cDNA	λ g t10		1.5 X 10 ⁶	0.5-3.4 kb (1.0 kb)	HL1084a

Figure 2. Region-Specific Human Brain Libraries.

These libraries are made from oligo(dT)-primed and randomly-primed cDNA from specific regions of the human brain. The tissues were obtained at autopsy from a disease-free 20 year old black male. The body was preserved at 4°C within one hour of death. Shown here are insert sizes of four random clear plaques from the \(\)\text{\text{gt11}} gyrus recti library, determined by using two \(\)\text{\text{gt11}} GeneAmp \(^{\text{TM}}\) PRI-MATE \(^{\text{M}}\) Amplimers (Cat No. 5412-1).

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.2 X 10 ^e	0.5-3.5 kb (1.1 kb)	HL1084b	
.4 X 10 ⁶	0.4-3.5 kb (1.0 kb)	HL1085a	
.0 X 10 ⁵	0.4-3.5 kb (1.1 kb)	HL1085b	
.2 X 10 ⁶	0.5-4.0 kb (1.2 kb)	HL1086a	
.2 X 1 ਹੈ	0.5 –3 .7 kb (1.1 kb)	HL10865	
.5 X 10 ⁶	0.6-4.0 kb (1.5 kb)	HL1087a	
.3 X 10 ⁶	0.6-4.0 kb (1.6 kb)	HL10875	
	.4 X 10 ⁶ .0 X 10 ⁶ .2 X 10 ⁶ .2 X 10 ⁶	(1.1 kb) .4 X 10 ⁶ 0.4-3.5kb (1.0 kb) .0 X 10 ⁶ 0.4-3.5kb (1.1 kb) .2 X 10 ⁶ 0.5-4.0kb (1.2 kb) .2 X 10 ⁶ 0.5-3.7kb (1.1 kb) .5 X 10 ⁶ 0.6-4.0kb (1.5 kb) .3 X 10 ⁶ 0.6-4.0kb	

L.E. c. ,	Ved	mRN tor Sour		# of Indep. Clones	insert Size Range (Average)	Catalog
Human Brain, Hippocampus cDNA	λgt1	0		1.0 X 10 ⁸	0.3–3.2kb (1.0kb)	HL1088a
Human Brain, Hippocampus cDNA	Àgi1	1		1.1 X 10 ⁶	0.3-3.2kb (0.8 kb)	HL:0885
arra Brain, Medulla	λ g (1)			1.3 X 10 ⁶	0.4–3.8kb (1.0 kb)	HL1089a
Human Brain, Medulia cONA	λ g :1 '			1.2 X 10 ⁶	0.4–3.7kb (1.1 kb)	HL1089b
Human Brain, Nucleus Accumbens cDNA	λgt10			1.0 X 10 ⁶	0.5–3.7 kb (1.2 kb)	HL1090a
Human Brain, Nucleus Thens cDNA	λ g (11			1.1 X 10 ⁶	0.6–3.8kb (1.1 kb)	HL1090b
ے سا Brain, Occipital Pole cDNA	λ g t10			1.8 X 10 ⁶	0.5–3.4 kb (1.1 kb)	HL1091a
Human Brain, Occipital Pole cDNA	λ g :11			1.6 X 10 ⁶	0.5–3.5kb (1.1 kb)	HL:0915
Human Brain, Putamen cDNA	λgt10			1.3 X 10 ⁶	0.4-3.3 kb (0.9 kb)	HL:092a
ಿ Brain, ಎಣಕಿಗ cDNA	λgt11			1.3 X 10 ⁸	0.4-3.4 kb (0.9 kb)	HL:092b
Human Brain. Substantia Nigra cDNA	λgt10			1.8 X 10 ⁶	0.6-4.0 kb (1.4 kb)	HL1093a
Human Brain, Substamia Nigra cDNA	lgt11			1.2 X 10 ⁶	0.6-4.0 kb (1.4 kb)	HL:093b
Human Brain, Superior aporal Gyrus cDNA	λgt10			1.0 X 10 ⁶	0.6-3.7kb (1.1 kb)	HL1094a
Human Brain, Superior Temporal Gyrus cDNA	λgt11			1.0 X 10 ⁸	0.4–3.7kb (1.2 kb)	HL1094b
Human Brain, Fetal, cDNA	λgt10	Fetal Brain to week old feto blood type:	ssue from 21 us. Mother's A+.	1.3 X 10 ⁶	0.6-4.0 kb (1.1 kb)	HL1065a
Human Brain, Fetal, DNA	λgt11	See above.		1.3 X 10 ⁶	0.7-4.0 kb (1.2 kb)	HL10655
Human Brain, Temporal Cortex cDNA	λgt11	Normal temp tissue, excise a tumor.	oral cortex od from around	7.3 X 10 ⁵	•	НL1003Ь

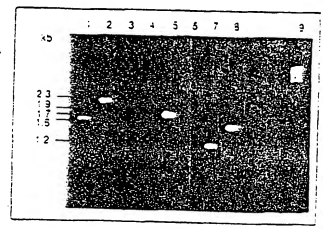
Library	Vecto	mRNA or Source	# of Indep. Clones	Insert Size Range (Average)	Catalog Number
Human Brain, Alzheimer's, cDNA	λgt11	Tissue around the hippocampus of a 70-year old diseased female, removed 5-6 hours after death.	8.1 × 10 ⁵	0.31-3.1 kb (1.1 kb)	HL1028b
Human Breast cDNA	λ g t10	Adult breast tissue excised during mastectomy, during the 8th month of pregnancy. Section showed well-differentiated tissue and lactational competence.	1.1 X 10 ⁶	0.6-3.3kb (1.1 kb)	HL:037a
Human Breast cDNA	اور 1	See above.	1.6 × 10 ⁶	0.5-3.4 kb (1.2 kb)	HL1037b
Human Breast, 5'-STRETCH cDNA	λgt10	See above.	1.4 X 10 ⁵	0.8 to>4.0 kb (1.5 kb)	HL1061a
Human Breast, 5'-STRETCH cDNA	λgt11	See above.	1.3 X 10 ⁶	0.8 to>4.0 kb (1.3 kb)	HL1061b
Human Breast, 5'-STRETCH cDNA	SWAJ-2	See above.	1.4 X 10 ⁶	0.8 to>4.0 kb (1.5 kb)	HL1061h
Human Breast Carcinoma, 5'-STRETCH cDNA	λ gt10	Breast carcinoma ZR-75-1.	1.3 X 10 ⁶	1.0 to>4.0 kb (1.7 kb)	HL1059a
Human Breast Carcinoma, 5'-STRETCH cDNA	λ gt11	Breast carcinoma ZR-75-1.	1.4 X 10 ⁶	1.0 to>4.0 kb (1.7 kb)	HL1059b
Human CEM T-Cell cDNA, randomly-primed + oligo(dT)-primed		CCRF-CEMT lymphoblastoid cell line (Foley et al., Cancer 18: 522 (1965).) Cell line is positive for CD4, CD5, T9 and T10, and negative for CD1, CD3, CD8, DR and TAC. Cell line is used for the propagation of HIV. ATCC # CCL 119.	1.6 X 10 ⁵	1.2 to >4.0 kb (1.7 kb)	HL1063f
Human CEM T-Cell cDNA, randomly-primed + oligo(dT)-primed	pBlue- script [†]	See above.	1.0 X 10 ⁶	1.1-4.6 kb (2.3 kb)	HL1063g
Human CML Spieen cDNA	1	Spleen of eight year old lemale with Chronic Myelogenous Leukemia.	1.4 X 10 ⁶	0.5–3.8 kb (1.1 kb)	HL1040b

<u>.</u> :	Vecto	mRNA or Source	# of Indep. Clones	Insert Size Range (Average)	Catalon
Human Colon cDNA	λ g t10	Normal tissues excised from around the colon cancer of a 53 year old male.	n 1.5 X 10 ⁶	0.6-3.2 kb (0.9 kb)	HL1034a
Human Colon cDNA	λ g 111	See above.	1.3 X 10 ⁶	0.6—3.5kb (1.1 kb)	HL1034b
₩ man Colon Tumor	λgt11 Not I/S/	Human T84 colonic tumor cell line.	1.3 X 10 ⁶	0.5–3.5kb (1.1 kb)	HL1079k
inuman Endotheli al cDNA	Agt1 1	Endothelial cells from umbilical cord veins. Cells were serially passaged and poly(A) isolated.	1.5 X 10 ⁶	0.34-3.4kb (0.9 kb)	HL10245
Human Endothelial cDNA, 5'-STRETCH	λ g t11	Endothelial cells cultured from human umbilical cord veins.	2.1 X 10 ⁶	0.6 to >4.0 kb (1.8 kb)	HL1070b
: _ an Epithelial cDNA	λgt11	Normal thymus epithelial cells, excised during open-heart surgery on a 3 year old Caucasian female.	1.4 X 10 ⁶	0.42–2.8kb (0.9 kb)	HL1025 5
Human Erythroleukemic cDNA		Human erythroleukemic K562 cells cultured in RPMI 1640 and treated with PMA as described. Siebert and Fukuda, J. Biol. Chem., 260: 640 (1985). Lozzio and Lozzio, Blood, 45: 321 (1975).	1.0 X 10 ⁶	0.5–3.4 kb (1.0 kb)	HL1032a
Human Eye cDNA	λgt10 /	Male/female eye pool.	1.1 X 10 ⁶	0.4–3.2kb (0.7 kb)	HL:047a
Human Eye cDNA	Agt11 p	dale/female eye pool.	1.5 X 10 ⁶	0.4-2.8kb (0.7kb)	HL1047b
Human Fallopian Tube 5'-STRETCH cDNA	λgt10 A	dult fallopian tube, nid-section.	1.3 X 10 ⁶	0.6 to>4.0 kb (1.5 kb)	HL1060a
Human Fallopian Tube 5'-STRETCH cDNA	λgt11 A	dult fallopian tube, iid-section.	1.2 X 10 ⁸	0.6 to>4.0 kb (1.5 kb)	HL1060b
Human Fibroblast, Lung	lgt11 Li	ung fibroblast cell line, IR-90. ATCC#CCL186.	2.2 X 10 ⁶	0.5~3.0kb (1.3 kb)	HL101:5
numan Fibroblast, Skin DNA	Agt10 C	ultured primary fibroblasts om a young male.	1.4 X 10 ⁶	0.5-3.6 kb (1.0 kb)	HL1052a
luman Fibroblast, Skin DNA	λgt11 Co	ultured primary fibroblasts om a young male.	1.1 X 10 ⁶		HL10525
					1 1/1

Library	Vector	mRNA Source	# of Indep. Clones	Insert Size Range (Average)	Catalog Number
Human Fibrosarcoma cDNA	λgt11	HT-1080 tumor cells. Rasheed et al., Cancer 33: 1027 (1974). ATCC# CCL121.	1.6 X 10 ⁶	1.0-3.4 kb (1.3 kb)	HL1048b
Human Genomic	EMBL3/ SP6/T7	Human Placenta DNA, Sau3A partial,	2.5 X 10 ⁶	8-21 kb (15 kb)	HL1067j
Human Glioma cDNA	λgt11	HS 683 from explant cultures of a glioma taken from the left temporal lobe. ATCC # HTB 138.	1.1 X 10 ⁶	0.4-3.1 kb (0.9 kb)	HL1049b
Human Hairy Cell Leukemia (Mo-B) cDNA	λgt10	Epstein-Barr virus-transformed B lymphoblast cell line. ATCC# CCL245.	1.4 X 10 ⁶	0.4–3.2kb (1.0 kb)	HL1043a
Human Heart cDNA	λgt11	Adult male heart, including some aorta region. Poly(A) RNA was slightly degraded as visualized on alkaline agarose gel.	1.27 X 10 ⁶	0.4-3.4 kb (0.9 kb)	HL1038b
Human HeLa Celi cDNA	λgt11	HeLa-derived D98-AH2 cells, HPRT- in phenotype. W.S. Szybalski et al., Natl. Cancer Institute Monograph, 7: 75 (1962).	1.3 X 10 ⁶	0.48-3.1 kb (0.86 kb)	HL1022b
Human Hepatoma cDNA	λgt11	Hepatoma cells, G2.	1.3 X 10 ⁶	0.5–3.7 kb (0.96 kb)	HL1015b
Human HUT-78 T-Cell 5'-STRETCH, cDNA, PMA-activated		Human T-cell line (<i>J. Exp. Med.</i> 154:1403 (1981).) Positive for IL-2. ATCC # TIB 161.	1.5 X 10 ⁶	1.0 to>4.0 kb (1.6 kb)	HL1068b
Human Keratinocyte cDNA	λgt11	Keratinocytes from adult epidermis.	1.7 X 10 ⁶	0.5-3.8 kb (1.1 kb)	HL1045b
Human Kidney cDNA	1	From a juvenile male whose kidney had been perfused for 24 hours prior to poly(A)* RNA isolation.	1.7 X 10 ⁶	0.5-3.2 kb (0.9 kb)	HL1033a
Human Kidney cDNA	λgτ11 :	See above.	1.4 X 10 ⁶	0.4–3.1 kb (0.9 kb)	HL1033b
Human Kidney, Fetal, cDNA, 5'-STRETCH	ſ	Fetal kidney tissues from a nixture of 20 week and 24 yeek old fetuses.	1.7 X 10 ⁶	0.8 to>4.0 kb (1.6 kb)	HL1071a

Liorary	Vector	mRNA r Sourca	# of Indep. Ciones	Insert Size Range (Average)	Calalog Number
Human Kidney, Fetal, cDNA, 3'-STRETCH	lgt11	Fetal kidney tissues from a mixture of 20 week and 24 week old fetuses.	1.7 X 10 ⁵	0.3 to>4 0 kb (1.5 kb)	HL10715
Human Laukooyta Ganomio	EMBL3	Laukocyte genomic DNA, maie.	3 8 X 10 ⁵	16ka	HU1008d
าก Laukocyte, Paripharal Blood, cDNA	रेदुर10	Peripharal blood leukocytes from aduit famale with Acute Promyelocytic Laukemia. HL60. S.J.Collins et al., Natura, 270: 347 (1977). ATCC#CCL240.	7.2 X 10 ⁵	0.69–3.5kb (0.38kb)	HL1020a

ere 3. Human Peripheral Blood Leukocyte cDNA Lety in λgt11. Cat. No. HL1062b. Seven randomly selected clear plaques were subjected to analysis using two λgt11 GanaAmp™ PCR^{††} PRI-MATE™ Amplimars (Cat No. 5412-1).



Human Leukocyte, Peripheral Blood, cDNA	λ g t11	See above.	8.4 X 10 ⁵	0.77 – 3.6kb (1.1 kb)	HL1020b
Human Leukocyte, ral Blood, SUNA, 5'-STRETCH	λ g t10	Adult male peripheral blood leukocytes.	1.7 X 10 ⁶	1 2 to>4.0 kb (1.6 kb)	HL:0623
Human Leukocyte. Peripheral Blood, cDNA, 5'-STRETCH	λ g t11	Adult male peripharal blood leukccytes.	1.8 X 10 ⁶	0.8 to>4.0 kb (1.6 kb)	HL1062b
Human Liver cDNA	10ءور	Normal adult liver, remaile.	1 5 × 10 ⁵	0.17-2.4kb (0.3kb)	HL:00'a
Tut - Liver cDNA	λgt11	Normal adult liver, female.	7 5 X 10 ⁵	0.2 –3 .0 kb (1.1 kb)	HL10015
Human Liver, Fetal, cDNA	Àgt11	Fetal liver, 1st trimester, male.	2.3 X 10 ⁵	0.14-2.3kb (1.1 kb)	HL:0055

Library	Vector	mRNA Source	# of Indep. Clones	insert Size Range (Average)	Catalog Number
Human Liver, Fetal, cDNA, 5'-STRETCH	λgt10	Liver tissue from 22 week old fetus. Mather's blood type: O+.	1.7 X 10 ⁶	1.2 to>4.0 kb (1.7 kb)	HL1064a
Human Liver, Fetal, 5'-STRETCH, cDNA	λ g t11	Liver tissue from 22 week old fetus. Mother's Blood type: O+.	1.5 X 10 ⁶	1.2 to>4.0 kb (1.6 kb)	HL10646
Human Lung cDNA	λ g t11	Normal adult lung tissue, excised during surgery.	8.2 X 10 ⁵	0.25–3.1 kb (1.2 kb)	HL1004b
Human Lung. 5'-STRETCH cDNA	λgt11	Adult lung tissue, including trachea and bronchioles.	1.4 X 10 ⁶	0.9 to>4.0 kb (1.5 kb)	HL1066b
Human Lung, Fetal, cDNA, 5'-STRETCH	λgt10	Fetal lung tissues from a mixture of 19 week and 21 week old fetuses.	1.4 X 10 ⁶	0.6 to>4.0 kb (1.5 kb)	HL1072a
Human Lung, Fetal, cDNA, 5'-STRETCH	λgt11	Fetal lung tissues from a mixture of 19 week and 21 week old fetuses.	1.3 X 10 ⁶	0.6 to>4.0 kb (1.6 kb)	HL1072b
Human Lung Fibroblast cDNA	λ gt11	Lung fibroblast cell line. IMR-90. ATCC#CCL186.	2.2 X 10 ⁵	0.5-3.0 kb (1.3 kb)	HL1011b
Human Lung Small Cell Carcinoma (NCI-H69) cDNA	λgt10	Lung Small Cell Carcinoma cell culture, NCI-H69.	1.4 X 10 ⁶	0.6–3.6 kb (0.93 kb)	HL1012a
Human Lung Small Cell Cardinoma (NCI-H69) CDNA	λgt11	Lung Small Cell Carcinoma cell culture, NCI-H69.	7.7 X 10 ⁵	0.4-2.8 kb (1.0 kb)	HL1012b
Human Lung Small Cell Carcinoma (NCI-H128) cDNA	λgt10	From NCI-H128 cell line derived from malignant pleural fluid of a 50 year old Black male. Gazder, et al., Cancer Research, 40: 3502 (1980).	1.0 X 10 ⁸	0.3–3.5 kb (0.85 kb)	HL1030a
Human Lung Smail Cell Carcinoma (NCI-H128) cDNA	Agt11	From NCI-H128 cell line derived from malignant pleural fluid of a 60 year old Black male. Gazder, et al., Cancer Research, 40: 3502 (1980).	1.3 X 10 ⁶	0.3-3.7 kb (0.92 kb)	HL1030b
Human Lung, WI-38, cDNA	λ g t1 1	WI-36 human diploid cell line, derived from normal Caucasian female lung. L. Hayflick, Exp. Cell Res., 25: 585 (1961).	1.2 X 10 ⁶	0 6-3.5 kb (1.1 kb)	HL10415

Libr	Vecto	mRNA or Source	# of Indep. Clones	Insert Siza Range (Average)	Catalog Number
Human Lymphocyte	λ g 110	Near confluent Raji cells, B lymphocytes. Pulvertaft, Lancet, 1: 238 (1964). ATCC# CCL86.	1.5 X 10 ⁶	0.25-3.2kb (0.87kb)	HL1002a
numan Mammary Gland		See Human Breast.			
· noma cDNA	λ g t11	Near confluent melanoma A2058 cells.	7.1 × 10 ⁵	0.6-4.0 kb (1.4 kb)	HL1023b
numan Monocyte, Peripheral Blood, cDNA	λ g τ1 1	90% human peripheral blood monocytes.	1.2 X 10 ⁶	0.5-3.4kb (1.1 kb)	HL10565
Human Monocyte, Peripheral Blood, cDNA.LPS-activated	λgt10	90% human peripheral blood monocytes, LPS-activated.	1.34 X 10 ⁶	0.4-3.5kb (1.2 kb)	HL:050a
nocyte, Blood, NA LPS-activated	λg:11	90% human peripheral blood monocytes, LPS-activated.	1.3 X 10 ⁶	0.5-3.7 kb (1.0 kb)	HL1050b
Human Monocyte, THP-1,cDNA	Agt10	THP-1 monocytes from 1 year old male with Acute Monocytic leukemia. Int. J. Cancer, 26: 171 (1980). Cancer Research, 42: 1530 (1982). ATCC# TIB202.	1.4 X 10 ⁶	0.55–3.0 kb (0.97 kb)	HL1021a
_ Monocyte, U-937, cDNA		U-937 cultured cells actively growing prior to poly(A) ⁺ RNA isolation.	1.5 X 10 ⁶	0.4-3.4 kb (0.95 kb)	HL1029a
Human Monocyte, U-937, cDNA	ِلُولِدًا 1 الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ الْجُولِدُ	See above.	1.2 X 10 ⁸	0.3-3.7 kb (0.95 kb)	HL1029b
Human Monocyte, PMA activated	e t	U-937 cells treated with 50ng/ml Phorbolmyristate acetate (PMA) for 3.5 days to achieve monocyte-like stage.	1.4 X 10 ⁶	0.6-3.8 kb (1.1 kb)	HL1035a
Human Monocyte, U-937, PMA activated cDNA	5 a to	U-937 cells treated with 50ng/ml Phorbolmyristate acetate (PMA) for 3.5 days o achieve monecyte-like stage.	1.2 X 10 ⁶	1.0-3.9 kb (1.3 kb)	HL1036b

Library	Vec	mRNA tor Source	# of Indep. Clones	insert Size Range (Average)	Catalon
Human Multiple Myeloma cDNA	λ g t11	Bone marrow obtained from a female with Multiple Myeloma, IM9. Ann.N.Y. Acad. Sci., 190: 221 (1972) PNAS 71: 84 (1974). J. Biol. Chem., 249: 1661 (1974). ATCC# CCL159.		0.52–3.5kb (1.1 kb)	HL1027b
Human Neuroblastoma cDNA	. \langle \langle \langle 10	Neuroblastoma cell line, Kelly.	1.05 X 10 ⁵	0.4-3.4kb (1.3 kb)	HL1007a
Human Osteosarcoma cDNA	λgiii	Osteosarcoma cell culture, MG-63. Antimicrob. Ag. Chemother., 12: 11 (1977). ATCC#CRL1427	1.8 X 10 ⁸	0.55-3.0 kb (0.97 kb)	HL10136
Human Ovary cDNA	λgt10	Normal ovary from 31 year old caucasian.	1.8 X 10 ⁶	0.6-4.0 kb (1.2 kb)	HL1098a
Human Ovary cDNA	λgt11	See above.	1.5 X 10 ⁶	0.6-4.0 kb (1.4 kb)	HL1098b
Human Pancreas cDNA	λgt11	Adult male pancreatic tissue.	. 1.4 X 10 ⁸	0.6–3.8 kb (1.2 kb)	HL1069b
Human Pancreas cDNA	SWAJ-	2 Adult male pancreatic tissue.	1.7 X 10 ⁶	0.6-3.5 kb (1.0 kb)	HL1069h
Human Pancreatic Carcinoma cDNA	λ g (10	Pancreatic cell line HS 766T. Ref.: J. Natl. Cancer Inst. 56: 843 (1976). ATCC# HTB134.	1.1 X 10 ⁶	0.5-3.4 kb (1.0 kb)	HL1057a
Human Pancreatic Carcinoma cDNA	λαι11	See above.	1.2 X 10 ⁸	0.5–3.4kb (1.0kb)	HL1057b
Human Peripheral Blood Leukocyte cDNA		See Human Leukocyte, Peripheral Blood cDNA.		(1.0 kg)	
duman Peripheral Blood Monocyte cDNA		See Human Monocyte, Peripheral Blood.		•	
luman Pltuitary cDNA		Tissue derived from a female with a gonadotropin-producing adenoma.	1.5 X 10 ⁸	0.6-4.0 kb (1.2 kb)	HL1096a
uman Pituitary cDNA	λgt11 :	See above.	1.7 X 10 ⁸	0.6-3.8kb (1.2 kb)	HL1096b

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Libr.		Vecto	mRNA or Source	# Inde Clos	of 9p. nes	insert S Rang (Averag	e Catalo
Human Pituitary cl		lgt10	Tissue derived from a female with a growth hormone-producing adenoma.	1.4 X	10 ⁶	0.5–3.8 k (1.2 kb)	b HI 1097
Human Pituliary cD. Inta cD		Agt11	See above.	1.3 X	106	0.6-4.0 ki (1.1 kb)	D HL10975
		λgt11	Placental tissue, 34 weel old.	ও 1.0 X 1	08	0.8–3.6 kb (1.8 kb)	
Human Placenta cDI 5'-STRETCH		λgt11	Placental tissue, 30 week old.	s 2.4 X 1	0 ⁶ 1	.2 to>4.0 l	с b HL1075b
Human Prostate cDN		G 110	Normal 65 year old prostatissues.	31e 1.5 X 10) ⁶	(1.8 kb) 0.6-3.7kb (1.2 kb)	HL1051a
State cDN, Human Retina cDNA		•	Normal 65 year old prostatissues.	ite 1.62 X 10	8	0.5–3.6 kb (1.1 kb)	HL10516
Human Retina cDNA			Adult retina.	1.6 X 10	6	0.5-3.5 kb (1.1 kb)	HL1055a
		111 /	Adult retina.	1.4 X 10 ⁶	•).5-3.4kb (1.0 kb)	HL:0556
Human Skin Fibroblast	هد	110 c	Cultured primary fibroblasts rom a young male.	1.4 X 10 ⁶	0	.5-3.6 kb (1.0 kb)	HL1052a
Human Skin Fibroblast	λgt	_	ultured primary fibroblasts om a young male.	1.1 X 10 ⁸	0.	5-3.5 kb 1.1 kb)	HL1052b
Human Spleen cDNA	λgt		ormal 25 year old male eleen.	1.4 X 10 ⁸	0.4	5-3.4 kb	HL1039a
iuman Spleen cDNA	λ g t1		ormai 25 year oid male leen.	1.3 X 10 ⁶	0.6	1.0 kb) 5-3.3 kb	HL1039b
luman Spieen, CML, DNA	\lambdagt1	Wit	leen of 8 year old female h Chronic Myelogenous ukemia.	1.4 X 10 ⁶	0.5	.0 kb) -3.8 kb -1 kb)	HL1040b
uman Stomach cDNA, •STRETCH	λgt1(mid a 62	ge of resectioned portion of stomach from year old male with nach carcinoma.	1.4 X 10 ⁵	0.6 to)>4.0 kb 4 kb)	HL1073a
man Stomach cDNA, STRETCH	λgt11		above.	1.7 X 10 ⁶	0.6 to	>4.0 kb	HL10735
man Submaxillary and cDNA	λgt11	Male biops	gland excised during Sy.	1.3 X 10 ⁶	0.5-) KD)	HL 1053b

CLONTECH Libraries

Library Human T-Cell cDN		Vector	mRNA Source		inde Clon	p.	nsert Ran Avera	ae .	Catalog
			T-Cell leukemic cell Jurkat, Schneider e J. Cancer, 19:621 (1 Yanagi et al., PNAS, 3430 (1985).	t al., Int.	1.1 X 1	6	0.58-3. (1.2 k	8 kb	HL1016a
Human T-Cell cDN		g(11)	See above.		1.1 X 10	•	. 58- 3.8	K D	HL1016b
Human T-Cell, CEM cDNA, randomly-pri + oligo(dT)-primed	Z.	ly (F (1 for an CE is t	CRF-CEMT mphoblastoid cell life color of the cell line is possible to the cell line is propagative to the propagative. ATCC # CCL 1	18: 522 ostive T10, CD3, oil line	1.6 X 10	⁵ 1.2	(1.2 kb to >4.((1.7 kb)) kb	HL1063f
HumanT-Cell,CEM cDNA, randomly-prime oligo(dT)-primed	pBli ed scri∣	ue- CC lym (Foi (196 for (and CD8 is us	RF-CEMT phoblastoid cell line by et al., Cancer 18 35).) Cell line is posi D4, CD5, T9 and T: negative for CD1, C. DR and TAC. Cell bd for the propagat V. ATCC # CCL 11	: 522 itive 10, D3, line	1.0 X 10 ⁶		-4.6kb .3 kb)	٠	HL1063g
Human T-Cell, HUT-78 5'-STRETCH, cDNA, PMA-activated	λgt11	Hum: Med.	In T-cell line (J. Exp. 154:1403 (1981).) Ve for IL-2 - ATCO #	1.5	5 X 10 ⁶	1.0 to> (1.6	4.0 kb kb)	HL	1068b
Human T-Cell, PEER 5'-STRETCH, cDNA Human T-Cell,	1gt10	ma, P		1.8	X 10 ⁶	1.0 to>4	I.O kb	HLI	078a
PHA Stimulated, cDNA	λgt10	blood o	ntrated T-Cell tion from peripheral if a healthy adult. mulated for 48 hour		X 10 ⁸	0.45 <u>-3</u> (0.95)	8kb	HL1	031a
uman T-Cell, HA Stimulated, cDNA uman Testis cDNA	λgt11	See abo		s. 1.0 X	108	0.7-3.3	kb	HL10	316
····~ CUIVA	λ g [] 1	healthy s	lesticle, excised urgery from a 50 year old. Not Ily induced.	1.0 X	10 ⁶	(1.4 kb 0.7–3.3 (1.2 kb	kb	HL10	

Library	Normal thumus		# of Indep.	113c11213		
Thymocyte CONA			Clones	(Average)	Catalog Number	
Human Thymus cDNA, 5'-STRETCH	λ g t11	old Caucasian female	1.7 X 10 ⁶	0.45-3.4kb (0.9 kb)	HL1026b	
S CONA,	SWA	1-2 See above.	1.5 X 10 ⁶	0.8 to>4.0 kb (1.6 kb)	HL1074b	
Human Thyroid Carcinoma cDNA	λ g (11	Adult thyrold carcinoma	1.5 X 10 ⁶	0.6 to>4.0 kb (1.5 kb)	HL1074h	
Human U251 cDNA. 5'-STRETCH	lgt10	ticsue, male. Human U251 Astrocytoma.	6.3 X 10 ⁵	0.65–2.7kb (1.2 kb)	HL1009b	
ਜਿਹਾਜ਼ਨ Wilms' Tumor ਂA	lgt11	Wilms' humar as "	1.5 X 10 ⁶	1.0 to>4.0 kb (1.8 kb)	HL1077a	
		from a 3 month old caucasian male. ATCC #	1.3 X 10 ⁵	0.6-3.6 kb (1.2 kb)	HL1044b	

Figure 4. Mouse Brain cDNA library in \(\lambda\) cat.

No. ML 1024a. Four randomly selected clear

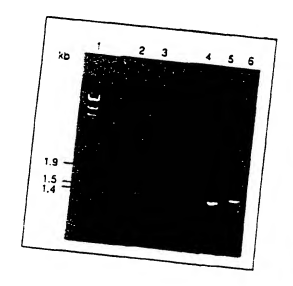
plaques were subjected to analysis using two \(\lambda\) gt10

\(\hat{Cat No. 5411-1}\).

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Molecular Cloning and Expression of the Human 55 kd Tumor Necrosis Factor Receptor പ്രാ

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Hans-Werner Lahm," Reiner Gentz,*
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Summary

Japan

Kamakura, Kanagawa

Two distinct receptors for tumor necrosis factor (TNF) of 55 and 75 kd are expressed at low levels by various cells. The 55 kd TNF receptor was purified from HL60 cells, and partial amino acid sequences were determined. Short degenerate sense and antisense oligonucleotide primers encoding the N- and C-terminal ends of a peptide of 22 amino acid residues were used to amplify a 66 bp cDNA fragment from HL60 RNA by reverse transcriptase-polymerase chain reaction. The cDNA fragment as a probe identified several overlapping clones in a human placenta cDNA library. The open reading frame of the cDNA predicts a 455 amino acid TNF receptor protein with leader, extracellular, transmembrane, and intracellular domains. When expressed in COS-1 cells or in a baculovirus system, the cDNA conferred TNF binding properties comparable to the native receptor. Surprisingly, the 55 kd TNF receptor shows a high degree of sequence homology to the NGF receptor extracellular domain.

Introduction

TNF-a and -β (or cachectin and lymphotoxin, respectively; jointly referred to as tumor necrosis factor [TNF]) are two cytokines with close functional and evolutionary relation (Nedwin et al., 1985): they compete for the same cellular binding sites (Aggarwal et al., 1985) and their genes are located in close proximity within the major histocompatibility complex in mouse and in man (Spies et al., 1986; Mueller et al., 1987). TNF was originally characterized as a factor with anti-tumor activity (Oid, 1985). However, the mechanism of tumor necrosis and regression remains poorly understood, even though TNF has been reported to exhibit direct cytotoxicity to many tumor cell lines in vitro (Matthews and Neale, 1987).

TNF has been found to possess a wide variety of biological activities. It exerts growth enhancing activity in fibro-

blasts, induces differentiation in human myeloid cell lines (for reviews, see Beutler and Cerami, 1987; Tracey et al., 1989), and has a crucial morphogenetic function in the induction and maintenance of granulomas in cell-mediated immunity (Kindler et al., 1989). Vascular endothelial cells are an important target for TNF. For example, TNF has been reported to induce expression of major histocompatibility HLA-A,B antigens or of adhesion-type molecules and to influence the morphology and proliferative activity of endothelial cell culture (Beutler and Cerami, 1987; Espevik et al., 1990). In other instances, TNF functions as mediator in immunologic and inflammatory responses. TNF concentrations in serum and cerebrospinal fluid correlate with the severity of inflammation in bacterial meningitis, with disease severity in malaria, or with mononuclear cell infiltration in acute phase graft-vs.-host disease (for reviews, see Beutler and Cerami, 1987; Tracey et al., 1989). TNF is one of the principal mediators of endotoxin in septic shock (Beutler et al., 1985; Rothstein and Schreiber, 1988; Michie et al., 1988).

Studies of intracellular signal transduction pathways revealed that TNF induces proteins that bind to κB -like enhancer elements and thus takes part in the control of NF- κB -inducible genes (Osborn et al., 1989; Lowenthal et al., 1989; Lenardo and Baltimore, 1989). The anti-viral activity of TNF at least in part is mediated by the interaction of NF- κB with a virus-inducible element in the β -interferon gene (Visvanathan and Goodbourn, 1989; Goldfeld and Maniatis, 1989). By an analogous mechanism, TNF appears to activate human immunodeficiency virus type I (Folks et al., 1989; Duh et al., 1989).

TNF-a in the crystal is packed in the form of a trimer of 17 kd monomeric units, and it is assumed that the trimer is the biologically active species (Hakoshima and Tomita, 1988; Eck et al., 1988; Jones et al., 1989). It cannot be excluded that a potential trivalency of the ligand, depending upon the surface density and lateral motility of receptors on various target cells, differentially affects receptor microclustering and thus has profound functional significance. In addition to the secreted 17 kd form, a 26 kd membrane-bound form of TNF is expressed in monocytes (Kriegler et al., 1988). It may function in intercellular contact or in paracrine fashion after posttranslational cleavage, resulting in the secretion of the 17 kd form.

The biological response to TNF is mediated by specific cell surface receptors. Several groups have reported studies on TNF receptors (Aggarwal et al., 1985; Creasy et al., 1987; Hirano et al., 1989; Hohmann et al., 1989; Kull et al., 1985; Niitsu et al., 1988; Smith and Baglioni, 1989; Tsujimoto et al., 1985). In TNF binding studies a single class of cell surface binding sites with a K_d in the pM range was identified on practically all of the cells analyzed. By cross-linking ¹²⁵I-TNF to the cell surface and 303-PAGE, however, two or even more bands specifically binding TNF were detected (Kull et al., 1985; Creasy et al., 1987; Stauber et al., 1988; Hohmann et al., 1989; Smith and Baglioni, 1989). Furthermore, soluble TNF

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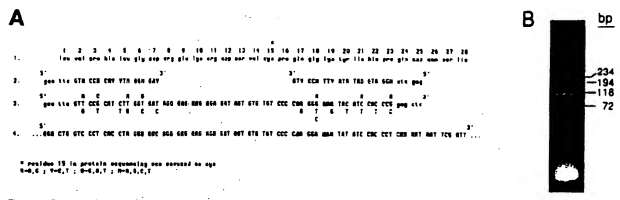


Figure 1. Partial Amino Acid Sequence of the 55 kd TNF Receptor and Gel Electrophoretic Analysis of the PCR Products

(A) Line 1: Sequence of the first 28 amino acids of the purified 55 kd TNF receptor. Residue 15 could not be asigned by protein sequencing but

(A) Line 1: Sequence of the first 28 amino acids of the purified 55 kd TNF receptor. Residue 15 could not be asigned by protein sequencing but was assumed to be cysteine. Line 2: Location and sequences of the sense and antisense primer mixtures. The primers were used for first-strand cDNA synthesis and PCR. Line 3: Nucleotide sequences of ten individual cDNAs cloned in pUC19 from the 78 bp band (see B). Line 4: Nucleotide sequence of the 55 kd TNF receptor cDNA in the region of the 28 N-terminal amino acids.

(8) The product of the PCR amplification, starting from first-strand rIL60 cDNA and using the sense and antisense primers defined in (A), fine 2, was electrophoresed through a 12% nondenaturing polyacrylamide gel and stained with ethidium bromide. The DNA of the 78 bp lower band was cloned in pUC19, and the nucleotide sequence of ten clones was determined (see A, line 3).

binding and inhibiting protein(s) of about 30 kd in human serum and urine were discovered (Seckinger et al., 1989; Olsson et al., 1989; Engelmann et al., 1989). Recently, another molecule, called Fas antigen, was described that does not bind TNF by itself but appears to be associated with TNF receptors and may play a role in signal transduction (Yonehara et al., 1989). In view of these complexities it is obvious that the various molecules involved need to be analyzed, preferably in cloning and transfection studies, to gain insight into their functional significance in the TNF/TNF receptor system.

In our previous work we have identified two cell surface molecules of 55 and 75 kd apparent molecular mass that specifically bind TNF-α and β with high affinity (Brockhaus et al., 1990; Hohmann et al., 1989). Specific monoclonal antibodies were raised to the purified molecules and were found to react exclusively with either the 55 or 75 kd molecule. The 55 kd molecule was found to be a functional TNF receptor, because anti–55 kd TNF receptor monoclonal antibodies elicited agonistic biological responses (Espevik et al., 1990). Here we report the molecular cloning and expression of the 55 kd TNF receptor and its relationship to the NGF receptor.

Results

Isolation of cDNAs Encoding the 55 kd TNF Receptor

The important steps in the isolation of 55 kd TNF receptor cDNA clones from a human placenta \(\lambda gt11 \) cDNA library are outlined in Figure 1. The 55 kd TNF receptor protein was purified by combined ligand- and immunoaffinity chromatography and reverse-phase HPLC from a HL60 cell lysate (Loetscher et al., unpublished data). The sequence of the first 28 N-terminal amino acids (and of internal peptides; see below) was determined by protein sequencing as snown in Figure 1A. Fully degenerate primer olioonucleotides were synthesized that encode amino

acids 17–23 (antisense primer) and 2–7 (sense primer) according to Figure 1A. Hexanucleotides containing the EcoRI and Sact endonuclease recognition sequences, respectively, were added to the 5' ends of the primer oligonucleotides. Total RNA was purified from HL60 cells, and first-strand cDNA was generated with reverse transcriptase, using the antisense oligonucleotide mixture as primer. Finally, the cDNA fragment encoding amino acid residues 2–23 was amplified in a polymerase chain reaction (PCR) using the combined sense and antisense primers and the first-strand HL60 cDNA.

The PCR product was analyzed on a native polyacrylamide gel and was shown to yield two sharp bands with the lower band running at the expected size of 78 bp (Figure 1B). This latter fragment was cloned into pUC19 by the attached restriction sites, and ten clones were sequenced. The sequence of the region between the two primer oligonucleotides was found to be invariant and to encode the correct amino acid sequence. In contrast, quite a number of different primers were used in the PCR (Figure 1A). The 78 bp cDNA fragment was used as a probe, and it led to the isolation of one clone containing the complete open reading frame and of four partial clones from the human placenta cDNA library.

The Predicted 55 kd TNF Receptor Is a Transmembrane Protein

The sequence of the 55 kd TNF receptor cDNA is shown in Figure 2A. The predicted amino acid sequence reveals a number of interesting features. The complete sequence consists of 455 amino acid residues with typical hydrophobic putative leader and transmembrane sequences as predicted by hydrophilicity analysis (Devereux et al., 1984). The single 21 amino acid transmembrane region separates an N-terminal, presumably extracellularly located domain of 182 amino acids including 24 cysteines, three potential N-linked glycosylation sites, and a C-terminal domain of 223 residues

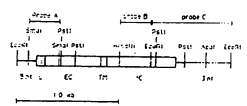
CAATTCGGGGGGGTTCAAGATCACTGGGACGCGGTGATCTCTATGCCCGAGTCTCAA CCCTCAACTGTCACCCCAAGGCACTTGGGACGTCCTGGACAGACCCAGTCCCGGGAAGCC -65 CCAGCACTGCCGCTGCCCACACTGCCCTGAGCCCAAATGGGGGGAGTCAGAGGCCCATAGCTG -28 . MetGlyLeuSerThrValProAspLeuLeuLeuProLeuValLeuLeuGluteu TCTGGCATGGGCCTCTCCACCGTGGCTGACCTGCTGCCCCTGGTGCTCCTGGACCTG -5 +1 LeuValGlyIleTyrProSerGlyValIleGlyLeuValProHisLeuGlyAspArgGluTTGGTGGGAATATACCCCTCAGGGGTTATTGGACTGGTCCCTCACCTAGGGGACAGGGG LysArgAspSerValCysProGlnGlyLysTyrileHisProGlnAsnAsnSerIleCys
AAGAGATAGTGTGTGTCCCAAGGAAAATATTCCACCCTCAAAATAATTCGATTCC 115 CysThrLysCysNislysGlyThrTyrLevTyrAsnAspCysProGlyProGlyGlnAspTGTACEAGTCCGACAAAGGAACCTACTTCTACAATGACTGTCCAGGCCCGGGCACGAT The Applyating Julyagius argly septhe the Alas erglus and is leurights accorded according accord CysleuSercysSerlysCyslerglysGlyMetGlyGlnValGluIleSerSerCysThr TGCCTCAGCTGCTCCAAATGCCCTAAATGCAGAATGCCTCAGGTGGAGATCTCTTCTTGCACA 295 ValkspärgäspänrvaldysGlyCysärgLysässGlnTyrärghisTyrärgSerGlu GTCCACCGGCACACTGTGTGTGTGCAGGAGAGCACCACTGGCATTATTGCACTGAA 90 355 AshleuPheGinCysPheAshlysSerLeuCysLeuAshGlyThrValHisLeuSerCys AACCTTTTCCAGTGCTTCAATTGCAGCCTCTGCTTGATTGCACCGTGCACCTTCCTGC CysValSerCysSerAsnCysLysLysSerLeuGluCysThrLysLeuCysLeuProGlnTGTGTCTCCTGTAGTAACTGTAACTAACAAAACCCTCCACGAAGTTGTGCCTACCCACGA 535 IleGiuAsnVallysGlyThrGluAspSerGlyThrThr<u>ValleufeuProleuVallle</u> 595 ATTGACAATGTTAAGGGCACTGAGGACTCAGGCACCACAGTGCTGTTGCCCCTGGTCATT Preparation of the Property of TrpLysSerLysLeuTyrSerIleValCysGlyLysSerThrProGlutysGluGlyGluTGGAAGTCCAAGCTCTACTCCATTGTTTGTGGGAATCGACACCTGAAAAGAGGGGGAG LeuGluGlyThrThrThrLySProLeuAlaProArnProSerPheSerProThrProJly CTTGAAGGAACTACTACTAGCCCCTAGCCCCAAACCTAGGTTCAGTCCCACTCCAGGC PheThrProThrleuGlyPheSerProValProSerSerThrPheThrSerSerSerThr TTCACCCCCACCCTGGCCTTCAGTCCGGTCCCACCTTCACCTCCACC TyrThr?rog.yAspCysProAsnPheAlaAlaProArgArgGluValAlaProProTyr TATACCCCCGGTGACTGTCCCAACTTTCCGCTCCCCCAGAGAGGTGCCACACCCTAT GlnGlyAlaAspProfleLeuAlaThrAlaLeuAlaSerAspProllePrcAsnProLeu CAGGGGGCTGACCCCATCCTTGCGACAGCCCTCCGACCCCATCCCCAACCCCCTT GInlystrpGluAspSerAlaHisLysProGlnSerLeuAspThrAspAspProAlathr CAGAAGTGGGAGGACACGCCCACAAGCCACAGGCCTAGACACTGATGACCCCGCGACG 1015 LeutyralavalvalGluasnvalFroProLeuArgTrpLysGlu2heValArgArgLeuCTGTACGCCGTGGTGGAGAACGTGCCCCCGTTGCGCTGGAACGAATTCGTGCGGCGCCCTA 1075 AlaGIntyrSerMetLeuklaThrTrpArgArgArgThrProArgArgGluAlaThrLeu GCGCAATACAGCATGCTGGCGACCTGGAGGCGGCGCGGGGCGAGGCCAGGCTG 1195 GluLeuleuGlyArg/alleuArgAspMetAspLeuIeuGlyCysleuGluAspIleGluGAGCTGCTGCGCACCGCTCCTCCGCAACATCGAC 1255 GluAlaLeuCysGlyProAlaAlaLeuProProAlaProSerLeuLeuArg GAGGGGCTTTGCGGGCCCGCCCCCCCCCCGCCCGGCCCLGGCTTCTCAGATGAGGCTGC GCCCCTGGGGGCAGCTCTAAGGACGTCCTGCGAGATGGCCTTCCAACCCCACTTTTTTC 410 1375 1495 1555 ACCCTATCCCTCATCCCCCTTTTGGGTGTCCTCACCAGCAAGGCTCCTCGGGGCCCCCTG 1675 1735 GTTTTGTTTTAAATCAATCATGTTACACTAATAGAAACTTGGCACTCCTGTGCCUTCTGCCCTGGACAAGCACAAGCACAAGCAGACAAGCTGAACTGTCCTAAGGCAGGGCGAGCACGGAACAATGG GCCCTTCAGCTGCAGCTGTGGACTTTTGTACATACACTAAAATTCTGAAGTTAAAAAAA MACCCGMATTC

Figure 2. The SS kd TNF Receptor Nucleotide and Predicted Amino Acid Sequences, and Schematic Representation of the 55 kd TNF Receptor cDNA Clone

(A) Amino acid numbering starts at the amino terminus Leu(+1); nucleotide numbering starts at the initiation codon. Amino acids 1-28 and 223-235 were also identified by sequencing the purified receptor protein. Residues 205-209 are in agreement with the receptor protein purified from human placenta (Loetscher et al., unpublished data). The putative transmembrane region is underlined. Potential N-linked glycosytation sites are indicated by asterisks. For the cysteine residue pattern, see Figure 6.

(B) Hybridization probes A (Sma!-Smal endonuclease-cut cDNA fragment), B (HindIII-EcoRI endonuclease-cut cDNA fragment), and C (EcoRI-EcoRI endonuclease-cut cDNA fragment) are indicated. The coding region of the cDNA is boxed and the putative leader (L), extracellular (EC), transmembrane (TM), intracellular (IC), and nontranslated (5'-nt, 3'-nt) regions are indicated.

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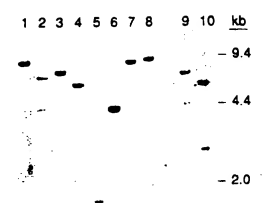


Figure 3 Genomic Southern Blot Analysis

Genomic DNA purified from HL60 cells was digested with restriction endonucleases (lane 1, EcoRi; lane 2, Hindill; lane 3, BamHI; lane 4, BamHI + EcoRI; lane 5, BamHI + Hindill; lane 6, Sspl; lane 7, Stul; lane 8, Apal; lane 9, Scal + Hindill; lane 10, Stul + Hindill), electro-phoresed through an agarose gel, and transferred to a nylon filter membrane. The filter was hybridized with probe 8 defined in Figure 28 as described in Experimental Procedures.

The reading frame in the C-terminal domain was confirmed by the sequence of an internal tryptic peptide fragment Ser(222)-Thr-Pro-Glu-Lys-Glu-Gly-Glu-Leu-Glu-Gly-Thr-Thr-Lys. An independent confirmation was provided by the sequence of a cyanogen bromide fragment of the 55 kd TNF receptor purified from human placenta starting with Tyr(205)-Arg-Tyr-Gln-Arg (Loetscher et al., unpublished data). Interestingly, the soluble TNF inhibitory protein from human serum and urine described by several other investigators (Seckinger et al., 1989; Olsson et al., 1989; Engelmann et al., 1989) appears to be a sciuble form of the 55 kd TNF receptor, because the 16–20 N-terminal amino acids described for the inhibitor are identical to a sequence of the 55 kd TNF receptor starting at aspartic acid(12).

A 29 amino acid leader sequence precedes the leucine identified by protein sequencing as amino terminus of the isolated mature protein leucine(+1); this predicts a signal peptide cleavage site that obeys the (-3, -1) rule (von Heijne, 1986), and thus leucine(+1) may represent the true N-terminus of the mature protein molecule. A more thorough analysis of a weight matrix of eukaryotic signal sequences, taking into account residues -13 to +1, reveals, however, a potentially more likely cleavage site at the Gly(-9)-lle(-8) peptide bond (von Heijne, 1986).

An unusual feature of the 55 kd TNF receptor sequence is the very high cysteine content in the extracellular domain (24 cysteines in a total of 182 residues). This explains why the tigand binding to the 55 kd TNF receptor is highly sensitive to reducing agents (Loetscher et al., unpublished data) and supports the hypothesis that the N-terminal sequence is the tigand binding and therefore extracellular domain. The high proline content of the receptor (34 residues) is another unusual feature expected from the results of amino acid composition analysis.

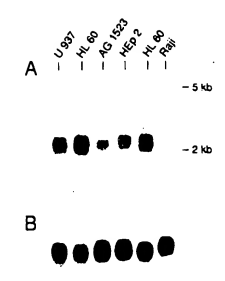


Figure 4. Northern Blot Analysis of Total RNA Purified from Various Cell Lines

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(A) RNA (12 µg per lane) was electrophoresed through a formaldehydecontaining agarcse gel and transferred to a nylon filter membrane. The filter was hybridized to probe 8 defined in Figure 28.
 (B) The same filter used in (A) was rehybridized after stripping to an actin probe as described in Experimental Procedures.

The 55 kd TNF Receptor Gene Is Unique

Genomic Southern blots of HL60 DNA were hybridized with the cDNA probes A, B, and C defined in Figure 28, which are located up- and downstream of the internal EcoRI site, respectively (see Figure 3 for hybridization with probe B; data for probes A and C are not shown). The two probes B and C identified two different bands in the EcoRI digest, whereas both probes identified the same bands in the BamHI, HindIII, Sspl, Stul, and Apal digests. Probe 8 hybridized with one additional band in Hindll digests, which is readily explained by a HindIII restriction site in an intron located between the Hindlil and EcoRI sites of the cDNA clone. Similarly, probe C hybridized with one additional band in Apal digests as a consequence of the Apal site in the 3' nontranslated region of the cDNA clone. Probes A and B hybridized to the same bands in the EcoRI, BamHI, and Apa; cigests. In HindIII digests, probe A identified a band of the same size as one that hybridized to probe B, but not to probe C. In Sspl and Stuldigests, probe A hybridized with bands that were different from those identified by both probes B and C. The structure of the 55 kd TNF receptor gene must be investigated in a more detailed analysis of genomic DNA; however, it appears from the present data that a single gene with at least three exons codes for the 55 kd TNF receptor.

The 55 kd TNF Receptor Is Expressed in Hi.60, U937, AG1523, and HEp2 Cells, but Not in Rail Cells

The pattern of 55 kd TNF receptor expression in HL60, U937, HEp2, AG1523, and Raji cells was studied by Northern blot analysis (Figure 4). A single mRNA species was

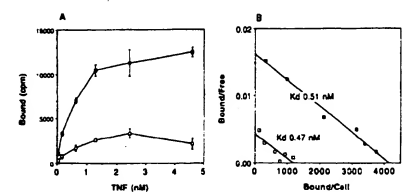


Figure 5. TNF Binding and Scatchard Analysis of COS-1 Cell Transfectants

(A) TNF binding. Monolayers of COS-1 cells transfected with the 1.3 kb EcoRI-EcoRI 55 kd TNF receptor cDNA as described in Experimental Procedures and nontransfected COS-1 cells in parallel were incubated with various concentrations of ¹²⁵I-TNF-a for 2 hr at 4°C and assayed for binding. Black squares, transfected COS cells; open squares, nontransfected control.

(B) Scatchard analysis of the binding data.

identified in HL60, U937, HEp2, and Ag1523 cells, whereas Rajl cells appeared to be practically devoid of 55 kd TNF receptor mRNA. These findings are in agreement with results of Western blot and flow cytometry studies using an anti-55 kd TNF receptor monoclonal antibody where HL60, U937, and HEp2 cells were clearly receptor positive, whereas no antigen was detected in Raji cells (Brockhaus et al., 1990). All three cDNA fragments (probes A, B, and C defined in Figure 2B) hybridized to the same band on the Northern blot. No evidence for the existence of a second mRNA species was obtained with probe A, which is specific for both the 55 kd TNF receptor and the TNF inhibitor sequences (Seckinger et al., 1989; Olsson et al., 1989; Engelmann et al., 1989).

Transfection and Expression of the 55 kd TNF Receptor cDNA Confers TNF Binding Activity

To establish that the 55 kc TNF receptor cDNA carries the complete minimum information required to confer specific TNF binding to a recipient cell, two independent expression systems were investigated. First, the 1.3 kb EcoRI-EcoRI fragment (Figure 2B) was cloned in a modified pXF3/ori vector, which contains the human cytomegalovirus immediate-early promoter and SV40 ori (Cullen, 1986). This construct was transfected into COS-1 cells (Gluzman, 1981) using lipofectin, and transient expression was measured (Felgner et al., 1987). Specific 125 TNF binding to the surface of the transfected cells was analyzed after 3 days in culture in the absence and presence of a 500fold excess of cold TNF. The binding data and Scatchard analysis are presented in Figure 5. The Kd of about 0.5 nM determined with the COS cell transfectants is comparable to the K_d of the native 55 kd TNF receptor as it is expressed almost exclusively on HEp2 cells (Hohmann et al., 1989). The endogenous TNF binding sites of COS cells have affinities comparable to those of the human 55 kd TNF receptor type. Furthermore, it was found that a monoclonal antibody, htr-9 (Brockhaus et al., 1990), directed to the human 55 kd TNF receptor extracellular domain inhibits TNF binding to the endogenous COS cell receptor almost completely. We assume, therefore, that the predominant endogenous COS cell TNF receptor is the homolog of the human 55 kd TNF receptor. Immunofluorescence studies using the anti-55 kd TNF receptor antibody (see above) show that the relatively low degree of expression of the 1.3 kb EcoRI-EcoRI fragment receptor construct is due to the fact that the ¹²⁵I-TNF binding measured in the assay (Figure 5) results from a small number of strongly positive transfectants, whereas the vast majority of cells is practically not stained.

Second, the 55 kd TNF receptor expression was investigated in a baculovirus expression system. The 1.3 kb EcoRI-EcoRI fragment was cloned in a modified pVL941 plasmid under the control of the polyhedrin promoter and introduced into the AcNP virus by homologous recombination (Luckow and Summers, 1988). When Sf9 cells were infected with the virus construct a highly significant and specific cell surface TNF binding was observed (see Table 1).

The 55 kd TNF Receptor and the NGF Receptor Have Highly Similar Extracellular Domains

A highly significant sequence similarity of the 55 kd TNF receptor with the nerve growth factor (NGF) receptor (Johnson et al., 1986; Radeke et al., 1987) was found. The alignment of both receptor sequences scored 13.6 standard deviations above the random score with the Mutation Data Matrix (Dayhoff et al., 1979). The similarity is most striking in the extracellular domains, where a block of 46 amino acids containing six regularly spaced cysteines is repeated four times in the two receptors (Figure 6). With suitable gapping, these domains containing 24 cysteines can be aligned, yielding 58 identities out of 169 possible matches (34% identity).

Table 1. TNF Binding with St9 Cells Infected with the 55 kd TNF Receptor cDNA AcNP Virus Construct

Cerls	Specific ¹²⁵ I-TNF Bound per 1 × 10 ⁶ Cells ^a
Noninfected Cells (Control)	60 cpm
Infected Cells ^b	1600 ± 330 cpm

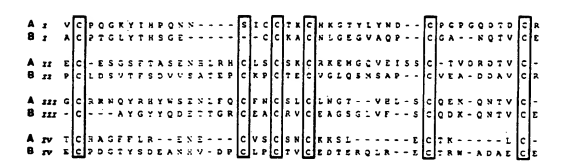
Virus construct, infection, and binding assay as described in Experimental Procedures.

Average and SD of four experiments.

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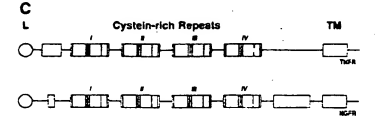


Figure 6. Homology between the 55 kd TNF Receptor and the NGF Receptor

(A) and (B) Four homologous cysteine-rich repeating elements in the extracellular domains of the 55 kd TNF receptor and the NGF receptor are shown (Johnson et al., 1986; Radeke et al., 1987). (A) 55 kd TNF receptor (I: residues 14–53; II: 54–96; III: 97–137; IV: 138–166). (B) Human NGF receptor (I: residues 4–36; II: 39–79; III: 81–118; IV: 121–180) (Johnson et al., 1986).

(C) Schematic comparison of the 55 kd TNF receptor and NGF receptor extracellular domain sequences.

Discussion

Our strategy for the molecular cloning of the 55 kd TNF receptor relieo to a large extent on PCR technology (Saiki et al., 1985). Despite its low abundancy, sufficient protein was purified to allow for the determination of 28 N-terminal and 15 internal amino acid residues. In preliminary experiments, the more conventional cloning approach using relatively short, fully degenerate or longer best-guess oligonucleotides as probes to screen cDNA libraries had proven technically difficult. An improved hybridization probe, therefore, was generated by PCR with degenerate primers and employed to identify the correct cDNA clone. A similar approach was used previously for the cloning of another, significantly more abundant gene (Lee et al., 1988).

The 55 kd TNF receptor has the typical structure of a membrane-spanning protein with a single transmembrane region separating intra- and extracellular domains. The calculated molecular mass of the mature protein is 47.5 kd. Because the purified receptor migrates with an apparent molecular mass of 55 kd on SDS-PAGE, post-translational modifications, most likely glycosylations, must account for the difference. Deglycosylation by N-glycanase treatment has been found to reduce the apparent molecular mass of the mature receptor by 5–10 kd (Hohmann et al., 1989; Loetscher et al., unpublished data), and therefore one or the other of the N-linked glycosylation sites appears to be utilized.

Recently, several lymphokine receptors, such as the erythropoietin receptor or the β chain of the IL-2, IL-4, and IL-6 receptors, have been shown to share extensive sequence homologies and to form a new interleukin receptor gene family (D'Andrea et al., 1989; Mosley et al., 1989;

Yamasaki et al., 1988). A distinctive element of this family is the Trp-Ser-X-Trp-Ser sequence motif found in the extracellular domain adjacent to the transmembrane region. The 55 kd TNF receptor lacks this motif and does not belong to this receptor family; rather, it is homologous in sequence to the NGF receptor, the sequence similarity being strongest in the extracellular domain, and belongs to the NGF/EGF/LDL receptor family.

In view of the potent bioactivity of TNF, a TNF inhibitor could have an important physiological role. The significance of the recently discovered TNF inhibitor peptide of human serum and urine (Olsson et al., 1989; Seckinger et al., 1989; Engelmann et al., 1989) is not yet understood. but it might function as a TNF sink. Our finding that the first 20 amino acids of the inhibitor match the 55 kd TNF receptor sequence starting at residue 12 of the mature receptor leaves little doubt that the inhibitor is a soluble fragment of the receptor molecule, probably containing most of the extracellular domain. Soluble forms of other. lymphokine receptors, such as the IL-2 receptor a chain, were reported previously (Rubin et al., 1985), and a specific mRNA encoding a soluble IL-4 receptor has been identified (Mosley et al., 1989). The genomic Southern blot analysis of the 55 kd TNF receptor provides no evidence for the existence of a distinct second gene that might encode the inhibitor. Furthermore, no evidence for the existence of a shorter mRNA species potentially encoding the inhibitor and created, e.g., by differential splicing, was found by Northern blot analysis of various cell lines. A specific TNF inhibitor mRNA, however, might be the result of a tissue-specific splicing event and thus need not be detected with the cell lines used in the present study. It is therefore possible that the TNF inhibitor is encoded by a tissue-specific differentially spliced transcript.

It seems more likely, however, that it is created by protectytic processing of the receptor molecule. With regard to the latter possibility, the finding of two amino termini, i.e., leucine(+1) in the receptor and aspartic acid(12) in the inhibitor (Seckinger et al., 1989; Olsson et al., 1989; Engelmann et al., 1989), and of a potential third signal peptide cleavage site predicted by sequence comparison, isoleucine(-8), is intriguing. It cannot be excluded that a primary translation product undergoes extensive posttranslational processing, which leads to the mature receptor and inhibitor molecules.

Several cell surface receptors have been found to consist of two or more peptide chains; for example, the IL-2 receptor system comprises at least a β and an inducible a chain, which function individually or as a complex (Hatekeyama et al., 1989). The TNF receptor system is no exception in this context: at present it comprises the 55 kd and 75 kd TNF binding molecules and the so-called Fas antigen (Yonehara et al., 1989). The 55 kg molecule, molecularly characterized in this study, is able to bind TNF and, upon binding of a specific antibody, to elicit biological responses in sensitive cells (Espevik et al., 1990). The availability of the 55 kd TNF receptor cDNA will allow investigation of TNF function and signal transduction in a more precise way than previously possible; it will also allow assessment of the roles of the 75 kd TNF binding protein and of other, potentially associated chains.

Experimental Procedures

Cell Lines and Growth Conditions

The human cell lines HL60 (ATCC CCL 240), U937 (ATCC CRL 1593). MEp2 (ATCC CCL 23), and Raji (ATCC CCL 86) and the dermal fibroblast line AG-1523 (Camden Cell Depository) were grown in RPMI 1540 or Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% horse or inactivated fetal calf serum. COS-1 cells (ATCC CRL 1650) were maintained in DMEM with 10% inactivated FCS. Sf9 insect cells (ATCC CRL 1711) were cultivated as described (Luckow and Summers, 1988).

Reagents

Recombinant human TNF-α purified from Escherichia coli was a gift from Drs. W. Hunziker, E. Hochuli, and B. Wipf (Hoffmann-LaRoche LTD, Basel). TNF-α was radioiodinated with Na ¹²⁵I (IMS40 Amersham) and Iodo-Gen (Pierce) to 0.3–1.0 x 10° cpm/mg as described (Fraker and Speck, 1978). Synthetic oligonucleotides were purchased from commercial sources.

Purification and Sequencing of the 55 kd TNF Receptor Protein

The 55 kd TNF receptor protein was purified to apparent homogeneity from HL60 cells by combined ligand- and immunoalfinity chromatography followed by reverse-phase HPLC on a Pharmacia ProRPC 5/2 column (Loetscher et al., unpublished data). Specific TNF binding activity of the purified protein was demonstrated in a solid phase assay (ligand blot). Briefly, proteins were separated by nonreducing SDS-PAGE and then electrophoretically transferred to a PVDF membrane (Matsudaira, 1987; Towbin et al., 1979). The membrane was blocked with 196 defatted milk powder and incubated with 5 ng/ml ¹²⁵I-TNF-α in the presence and absence of 5.0 µg/ml unlabeled TNF-α. TNF binding was detected by autoradiography.

An aliquot of the HPLC-purified receptor protein was subjected to automated Edman degradation, and the first 28 amino acids were determined. Additional sequence information was obtained after reduction and S-carboxymethylation of the protein (Jones, 1986), followed by cyanogen bromide cleavage (Tarr, 1986) and subsequent trypsin digestion in 200 mM NHL/HCO₃ (pH 8.0) (196 estimated motar railo trypsin,

added twice at 16 hr intervals at 37°C). The resulting peptide fragments were separated by reverse-phase HPLC on a Brownlee Aquapore RP 300 column (100 × 2.1 mm) and subjected to N-terminal sequence analysis on an Applied Biosystems 475A protein sequencer with online PTH-emino acid analyzer (Hewick et al., 1981).

To analyze similarities with other known sequences, sequence library searches and alignments were performed using the combined GenBank, National Biomedical Research Foundation, European Molecular Biology Laboratory, Protein Research Foundation, and Swiss-Prot data bases; computer programs were obtained from the Genetics Computer Group, the National Biomedical Research Foundation, and Hoffmann-LaRoche (Dayhoff et al., 1979; Devereux et al., 1984; Pearson and Lipman, 1968).

Polymerase Chain Reaction

First-strand cDNA was synthesized with total RNA purified from HL60 cells and a cDNA synthesis kit (Amersham) according to the instructions of the manufacturer, except that the degenerate antisense oligonucleolide mixture (Figure 1A) was used as primer for the reverse transcriptase. First-strand HL60 cONA in a mixture with the sense and antisense primer oligonucleotides (Figure 1A) was subjected to PCR using a Cetus GeneAmp kit and a Perkin-Elmer Thermocycler. The conditions were modified such that the primer concentrations were partially corrected for the level of their degeneracy. The PCR was run for 25 cycles (30 s at 94°C; 90 s at 55°C; 90 s at 72°C). An aliquol of the reaction was electrophoresed through a 12% nondenaturing polyacrylamide gel, bands were visualized by ethidium bromide staining and excised, and the DNA was recovered by electrophoretic transfer onto DEAE-cellulose paper as described (Maniatis et ai., 1982). To determine the nucleotide sequences, the PCR-amplified cDNA fragments were cloned in pUC19 vector. To obtain a screening probe the 78 bp fragment was labeled by PCR using 10 mM [a-32P]dCTP instead of 200 mM dCTP in the amplification reaction.

cDNA Library and Plaque Screening

A human placents cDNA fibrary ligated into \(\)gt11 vector was purchased from Clontech (1.0 \times 10^6 independent clones, 1.8 kb average insert size). The amplified library was plated, and duplicate plaque lift filters (GeneScreen Plus) were prepared according to standard protocols (Maniatis et al., 1982; Ausubel et al., 1989). The filters were hybridized to the denatured 78 bp cDNA probe (\(^{32}\text{P-labeled by PCR}\)) and washed as recommended by the manufacturer. Filters were exposed to X-Omat AR5 film with Cronex Li-Plus enhancer screen at -70°C overnight. Double positive clones were plaque purified, and the direct DNA was cloned in pUC and M13mp vectors according to standard protocols (Ausubel et al., 1989). DNA sequencing was performed with a Sequenase sequencing kit (US Biochemical).

Blot Transfer and Hybridization Protocols

For Northern blot analysis, total RNA purified from various cell lines was electrophoresed through 1.0% agarose gets containing formaldehyde as described (Maniatis et al., 1982). After electrophoresis gets were biotted by capillary transfer in 10× SSC to Zeta-Probe nylon membranes (BioRad). Southern blot analysis was performed by alkaline blotting DNA separated on agarose gets to Zeta-Probe nylon membranes as recommended by the manufacturer. cDNA probes were either labeled by random priming (DNA labeling kit; Boehringer Mannheim) or by PCR. Hybridization, washing, and stripping of the Zeta-Probe membranes was carried out in SDS buffer according to the instructions of the manufacturer. The final high stringency wash was in 40 m.M NaH₂PO₄, 1 mM EDTA, 1% SDS at 65°C performed twice for 30 min.

Cell Transfection, Expression, and Cellular Binding Assays

For transient COS-1 cell transfections (Gluzman, 1981), the 1.3 kb EcoRI-EcoRI fragment of the 55 kd *NF receptor cDNA was cloned in a modified pXF3/ori vector (Cullen, 1986), which contains the human cytomegalovirus immediate-early promoter and SV40 ori, and the construct at 1.8 μg/ml DNA was introduced into COS-1 cells by lipotectiniduced transfection (Felgner et al., 1987). After 2-3 days in culture the cells were detached with EOTA (Gibco) and tested for ¹²⁵-TNF-α binding. The cells were washed, resuspended at 2.8 × 10⁶ cells/ml, and incubated with various concentrations of ¹²⁵-TNF-α in the absence

and presence of a 500-fold excess of cold TNF- α for 2 hr at 4°C. The bound radioactivity was counted in a γ counter. Nonspecific binding was subtracted. Scatchard analysis was performed by the ligand algorithm (courtesy of P. J. Munson).

For the baculovirus expression system the 1.3 kb EcoRt–EcoRl fragment was cloned in a modified pVL941 plasmid under the control of the polyhedrin promoter and introduced into the AcNP virus by homologous recombination (Luckow and Summers, 1988). St9 insect cells (ATCC CRL 171) were infected, and after 3 days in culture the specific cell surface TNF-a binding was measured. The St9 cells were washed from the culture dish with a Pasteur pipet, resuspended at 5 × 10^s cells/mt in St9 medium containing 10 ng/ml ¹²⁸HTNF-a in the absence and presence of 5 µg/ml unlabeled TNF-a, and incubated for 2 hr on ice. The cells were washed with St8 medium, and the bound radioactivity was counted in a y counter.

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GenBank Accession Number

The accession number for the sequence reported in this paper is M33480.

Appendix I

Cell, Vol. 61, 361-370, April 20, 1990, Copyright © 1990 by Cell Press Best Avoil City Copyright

Molecular Cloning and Expression of a Receptor for Human Tumor Necrosis Factor

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Summary

A human tumor necrosis factor (TNF) binding protein from serum of cancer patients was purified to homogeneity and partially sequenced. Synthetic DNA probes based on amino acid sequence information . were used to isolate cDNA clones encoding a receptor for TNF. The TNF receptor (TNF-R) is a 415 amino acid polypeptide with a single membrane-spanning region. The extracellular cysteine-rich domain of the TNF-R is homologous to the nerve growth factor receptor and the B cell activation protein 8p50. Human embryonic kidney cells transfected with a TNF-R expression vector specifically bind both 125I-labeled and biotinylated TNF-α. Unlabeled TNF-α and TNF-β were equally effective at displacing the binding of labeled TNF- α to TNF-R expressing cells. Northern analysis indicates a single species of mRNA for the TNF-R in a variety of cell types. Therefore, the soluble TNF binding protein found in human serum is probably proteolytically derived from the TNF-R.

Introduction

Tumor necrosis factor-alpha (TNF-a) is a multipotent cytokine produced mainly by activated macrophages. TNF- α was originally identified as a tumoricidal protein effecting hemorrhagic necrosis of transplanted solid tumors in mice (Carswell et al., 1975) but has since been implicated in diverse biologic processes including inflammation and immunoregulation, antiviral defense, endotoxic shock, cachexia, angiogenesis, and mitogenesis (Goeddel et al., 1986; Beutter and Cerami, 1988; Old, 1988). The related cytokine lymphotoxin (TNF-β) is synthesized by activated lymphocytes and shares many of the biological activities of TNF-a (Goeddel et al., 1986).

The mechanisms through which the TNFs mediate their multiple activities are largely unknown, but like most polypeptide hormones, binding to specific cell surface receptors is an initial event. Stable trimers comprised of identical TNF-α polypeptides of 17,350 daltons bind to sites on a variety of cell types, with dissociation constants (V.)

ranging from 1.3 \times 10⁻⁹ M to 7.1 \times 10⁻¹¹ M (Aggarwal et al., 1985; Kull et al., 1985; Tsujimoto et al., 1985; Baglioni et al., 1985; Watanabe et al., 1986; Tsujimoto and Vileck, 1987; Stauber et al., 1988; Hohmann et al., 1989; Ding et al., 1989). While most investigators report a single class of cell surface binding sites, others report the presence of both high ($K_d = 2.6 \times 10^{-13} \text{ M}$) and low ($K_d = 1.5 \times 10^{-10}$ M) affinity sites on the same cell (Imamura et al., 1987). TNF- β and TNF- α have been shown to compete for binding to the same receptor on the human cervical carcinoma cell line ME-180 (Aggarwal et al., 1985) and the histiocytic lymphoma cell line U-937 (Stauber and Aggarwal, 1989). Estimates of the size of the TNF receptor (TNF-R) determined by affinity labeling studies range from 54 to 175 kd (Creasey et al., 1987; Stauber et al., 1988; Hohmann et al., 1989; Smith and Baglioni, 1989). A recent report suggests the existence of two major receptor types for TNF- α : a myeloid cell type receptor with a K_d of 7.1 imes 10⁻¹¹ M and an epithelial cell type receptor with a K_d of 3.2 imes 10⁻¹⁰ M. These two receptor types differ in size, glycosylation, and in their peptide maps (Hohmann et al., 1989).

In addition to the cell surface receptors for TNFs, several groups have identified soluble proteins in human urine (Peetre et al., 1988; Seckinger et al., 1988, 1989; Engelmann et al., 1989, 1990) and in the serum of numan cancer patients (Gatanaga et al., 1990) capable of specifically binding TNFs. In one instance two immunologically distinct TNF binding proteins (TNF BPs) were isolated from human urine (Engelmann et al., 1990). Antibodies raised against those two proteins (TBP I and TBP II) had an inhibitory effect on the binding of TNF-a to its cell surface receptor, suggesting a structural similarity between the cell surface TNF-R and the soluble TBPs.

Soluble cytokine binding proteins in biological fluids have been shown in some cases to represent "shed" forms of cell surface cytokine receptors (Rubin et al., 1985; Novick et al., 1989; Zupan et al., 1989). To ascertain whether this was the case for the TNF-R, we purified a soluble TNF 8P from human serum and isolated a corresponding cDNA by molecular cloning. This cDNA encodes a cell surface receptor for TNF that can presumably be processed to yield a soluble TNF BP.

Results

Purification and Characterization of TNF BP

A protein that inhibits the activity of both TNF- α and TNF- β has been detected in the serum of cancer patients but not healthy individuals (Gatanaga et al., 1990). This protein was purified by TNF- α affinity chromatography. Proteins eluted from the TNF-a affinity column were separated by reverse-phase HPLC, and several residues of N-terminal amino acid sequence were determined for each. Only one sequence was obtained, DSV(C/H)PQGKYIH, that did not correspond to a known serum protein. This protein, with an apparent M, of 28,000, showed N-terminal sequence

son et al., 1989; Engelmann et al., 1989, 1990). To obtain the amino acid sequence of internal peptides, the HPLC-purified TNF BP was subject to proteolysis using lysine C in the presence of SDS and the resulting proteolytic fragments were separated. The sequences of the two major eluting peaks were determined: GTYLYNDCPGFGODENE for PF I and EMGQVEiSSCTVDNDTVCG for PF II.

cDNA Cloning and Characterization Reveal a Receptor Structure

Two synthetic DNA probes were synthesized based on the amino acid sequence of PF I and PF II, using human codon bias (Lathe, 1985). The two probes were used to screen cDNA libraries made from placental tissue and from the promyelocytic cell line HL-60. Several positive clones were obtained from both libraries. The DNA sequence was determined for four overlapping cDNA clones from the HL-60 library. The composite sequence contained a single long open reading frame. The sequence of a 2.1 kb cDNA from the placental library was also determined and found to overlap the combined sequences of the HL-60 clones. The placental cDNA clone centains all of the presumed coding region as well as some of the 5' and 3' untranslated regions. The composite nucleotide sequence of the cDNAs and the deduced amino acid sequence of the predicted protein is shown in Figure 1A. There are three nucleotide differences between the placental and HL-60 clones (A at nucleotide position 75 in the placental vs. G in the HL-60, G vs. A at position 219, and G vs. A at position 1342), none of which results in an amino acid change. The open reading frame defines a protein of 455 amino acids starting at nucleotide position 182 and terminating at nucleotide position 1545.

The encoded protein exhibits a predicted domain structure typical of a cell surface receptor: the hydropathy profile indicates a signal peptide at the beginning of the protein and a potential transmembrane domain in the middle that separates the presumed extra- and intracellular domains (Figure 1B). The 11 amino acids designated +1 through 11 match the N-terminal amino acids of the soluble TNF BP isolated from serum. Therefore, we assigned Asp+1 as the N-terminal residue, although it is not known whether the N-terminus of the cell surface form of this molecule is the same as the N-terminus of the soluble form that was sequenced. Residues -40 to -10 are largely hydrophobic and probably serve as a signal peptide. Although the precise cleavage point is not known, the Gly (-12)-Leu (-11) peptide bond is a possible site (von Heijne, 1986). The mature protein may result from further proteolytic processing at the basic Lys (-2)-Arg (-1) dipeptide. Residues +1 through 172 probably constitute a cysteine-rich extracellular domain, with 24 cysteine residues and 3 potential sites for N-glycosylation (Asn-X-Ser/Thr) at residues 14, 105, and 111. In addition to the identity to the N-terminal sequence of soluble TNF BP, this domain contains sequences corresponding to the lysine C-generated proteolytic fragments PF I (residues 24-41) and PF II (residues 67-86). The 23 amino acid hydrophobic region in the middle of the molecule, which is flanked on its amino-terminal side by Thr 171 and on its carbone

terminal side by Arg 195, is characteristic of a transmembrane-spanning domain. The putative cytoplasmic domain would be comprised of the remaining 220 amino acids.

Cysteine Repeats in the Extracellular Domain of the TNFR

In the presumed mature extracellular domain of the predicted TNF-R protein, 24 of the 171 total amino acids are cysteines and the spacing of the cysteine residues is periodic. In contrast, the remaining 265 residues contain only 6 cysteines. Dot matrix analysis of these regions using the ALIGN score and the Dayhoff matrix (Dayhoff et al., 1978, 1983) reveals significant diagonal patterns of homology, indicating internal homologies (Figure 2A). Inspection of the extracellular domain sequence reveals it can be roughly divided into four related subdomains (Figure 2B). This 4-fold symmetry may represent duplication events of an ancestral subdomain in the evolution of the TNF-R.

TNF-R is Related to Nerve Growth Factor Receptor and the B Cell Activation Molecule Bp50

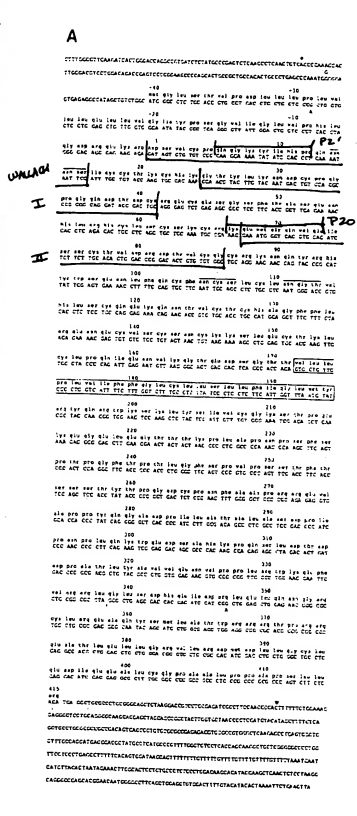
Cysteine-rich repeats have been detected in the extracellular domains of the EGF precursor and EGF receptor (Doolittle et al., 1984; Ullrich et al., 1984) and LDL receptor (Yarnamoto et al., 1984), but the TNF-R sequence reveals no significant amino acid homologies with these molecules. However, cysteine-rich extracellular domains have also been reported in the nerve growth factor receptor (NGF-R) (Johnson et al., 1986) and the B lymphocyte activation molecule Bp50 (CDw40) (Stamenkovic et al., 1989). These molecules have significant homology to the TNF-R in their extracellular domains (Figure 3). All three molecules share a similar set of cysteine-rich subdomains. Optimal alignment of the sequences for the extracellular domains of TNF-R, NGF-R, and Bp50 shows the close conservation of the cysteine residues and an overall identity of 29% between TNF-R and NGF-R and 24% between TNF-R and 8p50 in 167 residues (Figure 3). No significant homology is seen in the transmembrane or intracellular domains of these molecules.

TNF-R Transcript Is Expressed in a Variety of Cell Types

A panel of human cells and tissues was examined for the presence of TNF-R mRNA. Figure 4A shows expression of TNF-R mRNA in human term placenta and adult liver, as well as the breast carcinoma MB436, the nontumorigenic transformed breast epithelial cell line HBL100, the glioblastoma A172, and a primary squamous carcinoma, FG. In addition, cell lines resistant and sensitive to TNF cytotoxicity were tested for TNF-R mRNA. TNF-resistant cell lines (T-24 bladder carcinoma, A549 lung carcinoma) and TNF-sensitive cell lines (MCF-7 breast carcinoma and ME-180 cervical carcinoma) all exhibit TNF-R mRNA (Figure 4B).

The increased expression of TNF-Rs on colls after treatment with interferon-y has been reported (Aggarwal et al., 1985). To test whether the levels of transcript for the TNF-R

Cioning of a Human TNF Receptor



 $-\beta$, we treated A549 cells with these cytokines for 24 hr prior to harvest of the mRNA. Levels of TNF-R mRNA are relatively constant after treatment with these cytokines alone or in combination (Figure 4C). Thus, if interferon-y or TNF regulation of receptor does occur in A549 cells, it does not appear to be transcriptional or it is mediated

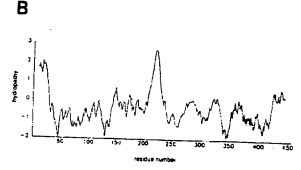


Figure 1, Human TNF-R cDNA and Amino Acid Sequences and Hydropathy Plot

(A) Composite nucleotide and deduced amino acid sequence of TNF-R cONA. Amino acids are numbered in the sequence. The composite of HL-60 clones begins at the first nucleotide and ends at nucleotide 1607 (f v). The placental clone begins at nucleotide 67 (f f O) and ends at the last nucleotide. Three nucleotide differences at positions 75, 217, and 1342 are noted, with the top nucleotide representing that found in the placental clone and the bottom representing the HL-60 sequence. Aspartic acid (Asp) is the initial residue in the N-terminal sequence obtained from native TNF BP in human serum and is labeled as position 1. Amino acids -40 to -1 comprise presumed signal peptide and proteolytically processed N-terminal residues. Overlines represent three potential N-linked glycosylation sites (residues 14, 105, and 111). The predicted transmembrane region (residues 172-194) is boxed. (9) Hydropathy profile of protein sequence in (A).

. through another class of receptor. We do not know whether TNF-R mRNA is affected by these cytokines in other cell

TNF-R message was also found in several hematopoietic cell lines: The cultured T cell lines CEM, HSB, and HuT 78, the functional cultinoir T cell line Distract the

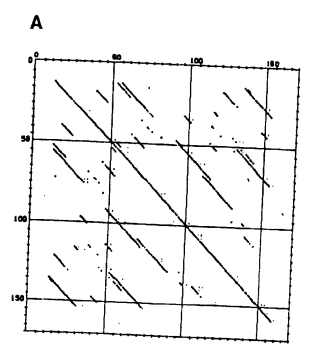


Figure 2. Internal Repeats in the TNF-R

- (A) Dot matrix piot of internal homologies in the TNF-R extracellular domains. Dots are placed where Dayhoff mutation matrix alignment scores >20 are obtained (Dayhoff et al., 1983). The plot is necessarily symmetrical around the diagonal line of identity; other lines at 45° angles that are off of the main diagonal represent areas of internal homology. Numbers denote amino acid positions in the predicted mature extracellular domain.
- (B) Internal cysteine repeats in the TNF-R extracellular domain. Alignment of amino acid sequences in the extracellular portion of the TNF-R represent four internal subdomains. Identical residues are boxed; amino acids are numbered in the left margin.



erythroleukemia line K562 have a single detectable species of TNF-R mRNA (data not shown). The cultured B cell line RPMI-1788, but not the EBV-transformed B cell lines JY, LB, and BOC, have TNF-R mRNA (data not shown). Uninduced U-937 cells were found to have relatively high levels of TNF-R mRNA, and uninduced HL-60 cells contained markedly less (data not shown). Thus, message for the cloned TNF-R seems widely though not ubiquitously expressed. In all cells that express TNF-R, a single species of mRNA of about 3.0 kb is detected. This suggests

that the cDNA clones we obtained do not represent complete copies of the TNF-R mRNA; some nucleotide sequences in the untranslated regions are missing.

Transfection and Expression of the Human TNF-R CDNA

The 2.1 kb placental cDNA clone was inserted into the mammalian expression vector pRK5. This cDNA starts at nucleotide position 64 (Figure 1A), with the initiating me-

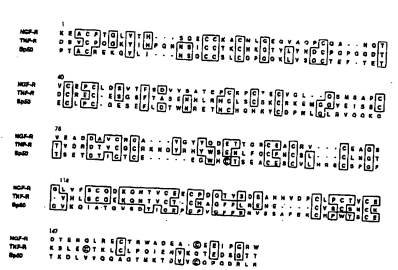


Figure 3. Homology in the Extracellular Domains of NGF-R, TNF-R, and 8p50

Optimized alignment of the protein sequence of the extracellular domains of NGF-R, TNF-R, and 8p50 (CDw40) is shown with gaps introduced to optimize matches. Identical amino acids are boxed. Cysteine residues not conserved among all three sequences are circled. Residues are numbered in relation to their position in TNF-R.

Cloning of a Human TNF Receptor 365

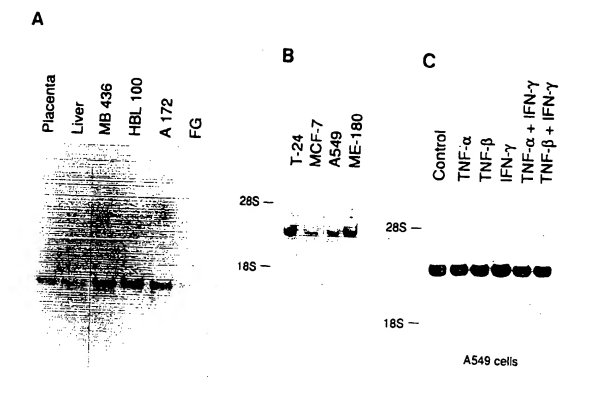


Figure 4. Northern Blot Analysis of TNF-R mRNA

(A) Northern analysis of TNF-R mRNA in placenta, liver, and transformed cell lines. Northern blot analysis of 4 μg of poly(A)* RNA from normal human term placental and adult liver tissues, the human transformed cell lines MB43€, HBL100, and A172, and the primary squamous carcinoma FG. (B) Northern blot analysis of both TNF-resistant and TNF-sensitive cell lines. TNF-resistant cell lines: A549 and T-24. TNF-sensitive cell lines: MCF-7 and ME-180. There is 3 μg of poly(A)* RNA per lane. The positions of ribosomal RNA bands are denoted as 28S and 18S. (C) Levels of TNF-R mRNA in A549 cells after IFN-γ or TNF treatment. There is 7 μg of poly(A)* RNA per lane from A549 cells after treatment with either TNF-α, TNF-β, or interferon γ, or both (0.1 μg/ml each) for 24 hr prior to harvest of mRNA. Control lane: untreated cells.

thionine 118 bp downstream (position 182, Figure 1A). The cDNA is under the transcriptional control of the cytomegalovirus immediate-early promoter and is followed by the SV40 termination and polyadenylation signals. The TSA 201 cell line, a subclone of the human embryonic kidney cell line 29,3s (Graham et al., 1977), which constitutively expresses large T antigen, was selected for transient transfection experiments because of its high transfection efficiency and low numbers of endogenous TNF-Rs. Following transfection with the TNF-R expression construct, cells were tested for the ability to bind specifically either biotinylated or ¹²⁵I-labeled TNF-a.

An increase in the relative capacity of pRKTNF-R-transfected cells to bind biotinylated TNF- α can be seen using fluorescence-activated cell sorting (FACS). Mock-transfected TSA 201 cells display a low level of binding of biotinylated TNF- α , presumably to endogenous receptors (Figures 5A and 5B). The levels of TNF- α binding are substantially increased on cells transfected with the TNF-R construct, as shown by the shift to the right of the fluorescence histogram (Figure 5F). The intensity of fluorescent staining is reduced to background levels in both populations by preincubation of cells with either nonbiotinylated TNF- α -or TNF- β (Figures 5C, 5D, 5G, and 5H), demonstrating that the observed binding is specific.

A saturation isotherm for the specific binding of 125]. TNF-a was performed by sequential dilution of the spe-

citic activity of the radioligand with unlabeled TNF- $\!\alpha\!$ at concentrations ranging from 67 pM to 33 nM (Figure 6A). The specific binding of 125 ITNF- α is saturable, and the Scatchard analysis of these data using nonlinear leastsquared regression reveals two binding sites (p <0.05; Figure 6A, inset) with high ($K_d = 0.56 \text{ nM}$) and low ($K_d = 0.56 \text{ nM}$) and $K_d = 0.56 \text{ nM}$) and $K_d = 0.56 \text{ nM}$ 19.6 nM) affinity. The number of binding sites for these two receptor subtypes on the transiently expressing TSA 201 cells is \sim 50,000 and 630,000 sites per cell, respectively. 125]-TNF-a binding to mock-transfected control cells was at least 10-fold lower than pRK-TNF-R-transfected cells, suggesting low numbers of endogenous receptors. Increasing concentrations of unlabeled TNF-B cause a dose-dependent decrease in specific 1251-TNF-a binding in a manner very similar to that that seen with unlabeled TNF-a (Figure 6B). This suggests that the TNF-R expressed on the transfected cells recognizes TNF- α and TNF-β with approximately equal affinity.

Discussion

The data presented here describe the cloning and expression of a receptor for human TNF-o and -\(\beta\). The deduced amino acid sequence of the TNF-R reveals a structure typical of cell surface receptors for polypeptides: it contains a signal peptide and extracellular, transmembrane, and intracellular regions. There is a significant degree of homol-

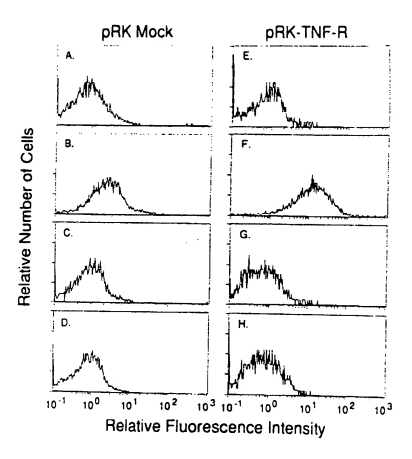


Figure 5. FACS Analysis of pRK-TNF-R-Transfected TSA 201 Cells

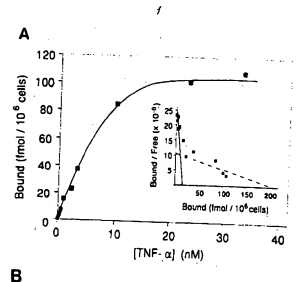
The left column contains histograms from mock pRK5-transfected cells; the right column contains histograms from pRK-TNF-R-transfected cells. Cells were stained with PE-conjugated streptavidin after pretreatment with or without biotinylated TNF-α (b-TNF-α). (A) and (E), PE-streptavidin without b-TNF-α; (B) and (F), PE-streptavidin plus b-TNF-α; (C) and (G), preincubation of the cells with a 5-fold excess of unlabeled TNF-α prior to staining with PE-streptavidin plus b-TNF-α; (D) and (G), preincubation with a 2-fold excess of unlabeled TNF-β prior to staining with PE-streptavidin plus b-TNF-α.

ogy between the TNF-R and the NGF-R, as well as the B lymphocyte activation molecule Bp50, suggesting that they comprise a family of related growth factor receptors. The four 40 amino acid repeats of the extracellular portions of these receptors may form relatively independent folded subdomains, each tightly cross-linked by disulfide bonds. Because these cysteine-rich domains form virtually all of the extracellular portion of the TNF-R, it is likely that they contain the ligand binding site(s), although it is not immediately clear how the biologically active TNF trimer binds the receptor. In this regard it is interesting to note that the net change of the extracellular domain of the TNF-R is positive, while that of its ligand is negative. The converse is true for NGF-R and its ligand, where the extracellular domain is strongly acidic and the ligand is basic. This would suggest that electrostatic attractions play an important role in the interactions of these related receptor molecules with their respective ligands. One possibility is that the cysteine repeat units form a general structural framework that, depending upon specific amino acid substitutions, could be varied to accommodate a variety of ligands.

The mechanisms of signal transduction by TNF and its receptor are obscure. The intracellular domain, though large enough to possess an enzymatic activity or to interact with other proteins that may mediate signal transduction, has no apparent sequence homology with any proteins in the available data bases. The cytoplasmic domain is rich in serine, threonine, and tyrosine but shows no homology to the catalytic domain of the tyrosine or ser-

ine/threonine-specific protein kinases. However, there is sequence similarity to the canonical phosphorylation sites (Ser/Thr-X-Arg/Lys) that can be acted upon by protein kinase C (Woodget et al., 1986) at amino acid positions 223, 366, and 371. Also present are a potential cyclic nucleotide-dependent protein kinase phosphorylation site at amino acid 368 (Feramisco et al., 1980) and a consensus tyrosine kinase phosphorylation site at residue 354 (Patchinsky et al., 1982). The binding of TNF to its receptor has been shown to increase both GTP binding and GTPase activity in HL-60 membrane preparations, leading to the suggestion that a GTP binding protein might couple TNF-induced signaling to biological effects (Imamura et al., 1988). However, the TNF-R has no homology to other receptors that are known to interact with G proteins.

Internalization of the native TNF-R in the absence of ligand has been shown (Smith et al., 1990; Ding et al., 1989) and shown to occur more rapidly in the presence of ligand (Smith et al., 1990; Imamura et al., 1987). While a majority of reports indicates the degradation of receptor after internalization (Watanabe et al., 1988; Smith et al., 1990), one suggests that the receptor is continuously recycled (Vuk-Pavlovic and Kovach, 1989). These differences could represent cell type—specific differences or differences in processing for distinct types of TNF-R. Regions of the intracellular domain of the cloned TNF-R reported here are extremely rich in proline, glutamic acid, serine, and threonine. The presence of these so-called PEST se-



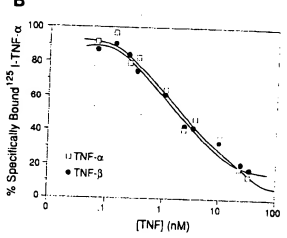


Figure 6. Binding Characteristics of Recombinant Human TNF-R Expressed in TSA 201 Cells

(A) Saturation isotherm of specific binding of ¹²⁵I-labeled TNF-α on transfected cells. Replicate samples of TSA 201 cells transiently expressing pRK-TNF-R were incubated with ¹²⁵I-TNF-α (16.7 pM) alone or in the presence of increasing amounts of unlabeled TNF-α, and the specific binding of ¹²⁵I-TNF-α was determined at each concentration. The inset presents the data transformed by Scatchard analysis. These results were from a single experiment that had been repeated three times with either triplicate or duplicate determinations.

(B) Displacement curves showing inhibition of the specific binding of 1251-TNF- α by unlabeled TNF- α (\square) or TNF- β (\blacksquare). The experiment was performed as described in (A), with 1251-TNF- α displaced with increasing amounts of TNF- α or - β .

lar protein degradation (Rogers et al., 1986). This leads to the speculation that the receptor encoded by this gene has a high turnover rate in its native state. This speculation is consistent with our observations that A549 cells have relatively high levels of mRNA for the TNF-R yet relatively low numbers of cell surface receptors (unpublished data).

It is not clear that the effector functions of TNF are achieved by transduction of a signal via the receptor after ligand binding. A direct intracellular role for TNF in cytotoxicity has been proposed based on TNF microinjection experiments (Smith et al., 1990), although using other cell lines we have not been able to demonstrate activity for microinjected TNF (D. Pennica and D. V. G., unpublished data). If internalization of the figand is the important step

in the biochemical action of TNF, the receptor may play no more a role than transporting TNF to the inside of the cell. However, the fact that antibodies generated against soluble TNF BPs will cross-react with cell surface molecules and act as TNF agonists suggests that TNF-R signal induction can occur without internalization of the TNF (Engelmann et al., 1990).

Shedding the extracellular domain of the TNF-R might be used as a protective mechanism by cells to avoid the cytotoxic effects of TNF. The presence of soluble TNF BP in the serum of cancer patients may represent a mechanism by which tumors evade host anti-tumor defenses by modulating systemic levels of TNF. Soluble forms of the receptor for IL-2 are released from activated human lymphoid cells (Rubin et al., 1985), and its levels are found increased in bodily fluids in disease states (Marcon et al., 1988). Soluble receptors for IL-6 and IFN-y have recently been detected in human urine (Novick et al., 1989), soluble truncated forms of the NGF receptor have been seen in human urine and amniotic fluid (Zupan et al., 1989), and a cDNA that encodes a soluble form of murine IL-4 has been reported (Mosley et al., 1989). The prevalence of such soluble receptors suggests a normal regulatory role for these molecules. Their mode of action could be to limit the amount of available cytokine by binding it in solution, thus preventing the cytokine from reaching its cell surface target. TBP I, the soluble TNF-BP that shares N-terminal homology to the TNF-R reported here, binds TNF-a with greater affinity than TNF-B (Engelmann et al., 1990). Our studies show that TNF-R binds TNF-α and -β with approximately equal affinities. This suggests that solubilization of the extracellular domain of the receptor may induce a change in the relative affinities of the binding component for TNF-α and TNF-β.

Most if not all mammalian cells appear to have receptors for TNF. However, the number of TNF-Rs per cell is relatively low. In the 5' untranslated region of TNF-R mRNA two short open reading frames are seen, one of 43 codons and one of 3 codons. Such short open reading frames are not uncommon in growth factor receptor RNAs. They are found 5' of the main open reading frame in the GM-CSF R (Gearing et al., 1989), the human IL-6 R (Yamasaki et al., 1988), the murine IL-1 R (Sims et al., 1988), and the human IL-2 R (Nikaido et al., 1984; Hatekeyama et al., 1989). It has been postulated that these short open reading frames might act, if translated, to dampen the translation of the main receptor coding regions (Gearing et al., 1989). Such a mechanism might partly explain the low numbers of these receptors, in relation to the levels of mRNA, on normal cell types.

Interpretation of published data from affinity labeling cross-linking studies is made difficult because cross-linked ligand is itself resolved into monomers, dimers, and trimers by SDS-PAGE, but estimates of the size of cell surface TNF-Rs is generally between 55 and 138 kd (Creasey et al., 1987; Stauber et al., 1983; Hohmann et al., 1989). Receptors of considerably larger size, up to 310 kd, have also been reported (Smith and Baglioni, 1989), but these forms might reflect receptors cross-linked to associated regulatory proteins, or even complexes of cross-

linked receptors. Association with other proteins or "adaptor subunits" has been shown for other receptors, most notably IL-2 R (Teshigawara et al., 1987; Hatekeyama et 🤛 al., 1989) and IL-6 R (Yamasaki et al., 1988). In addition, NGF-R is thought to require association with another molecule for high-affinity binding of its ligand (Radeke et al., 1987). Association with an as yet unknown protein may be required for the TNFR reported here to bind ligand with high affinity. One possible explanation for the two binding sites observed in the cells transfected with pRK-TNF-R is that a small number of the transiently expressed receptor proteins interacts with an endogenous protein present in a limited quantity within the TSA 201 cell. This subpopulation of receptors might then bind 1251-TNF-a with a higher relative affinity ($K_d = 0.68$ nM), while the bulk of the expressed receptors (92% based on B_{max} predictions) binds ligand with ~30-fold lower affinity.

The relationship between the TNF-R described here and other potential TNF-Rs is not clear. The predicted 415 amino acid molecule we identified by cDNA cloning encodes a protein with a predicted Mr of 50,578. Since there are three potential N-linked glycosylation sites, and since protein bicchemical studies consistently reveal carbohydrate content on the TNF-R molecules analyzed, it is likely that the apparent Mr of the TNF-R reported here is greater than 50,000 in its native state. The recent report of two immunologically distinct forms of TNF BPs in urine, one of which has N-terminal sequence homology to the sequence reported here (Engelmann et al., 1990), argues for the existence of at least two cell surface TNF-Rs that shed their extracellular domains. This is consistent with the report of two major types of TNF-R on different cell types (Hohmann et al., 1989), suggesting that the molecute reported probably represents one of the two types of cell surface TNF-R. The machanisms by which the extracellular domains of the TNF-R are shed are not known. Soluble IL-4 receptor is thought to be the result of alternative splicing of IL-4 mRNA (Mosley et al., 1989). In contrast, the presence of only one detected species of mRNA for this TNF-R suggests that the soluble form is generated proteolytically by cleavage of the extracellular domain from the cell surface receptor. Soluble TNF BP may be resistant to subsequent proteolysis after release as a result of a compact disulfide-bonded structure. The physiological significance of this process remains unknown. The availability of this and other cloned TNF-R cDNAs should allow for the resolution of these issues as well as the elucidation of many of the complexities of the multiple activities of TNF.

Experimental Procedures

Purification of Serum TNF BP and Amino Acid Sequencing of Proteolytic Fragmenta

TNF-a affinity chromatography fractions of serum proteins from human cancer patients were shown to inhibit the activity of TNF-a and TNF-B (Gatanaga et al., 1990). These fractions were electrophoresed on SDS-PAGE and found to contain several components. Samples from the affinity column were loaded directly onto a small glass column (1.5 mm \times 50 mm) packed with 15 μ m of \bigcirc -18 packing material (J. T. Baker) and eluted on a HP 1090 HPLC with a linear gradient of 1%-60% acetonitrile in 0.1% TFA and water at a flow rate of 0.2 ml/min. Eluted

peaks were sequenced directly on a prototype sequencer (U.S. patent number EP0257735), and sequences obtained were compared to the protein sequences in the available data bases using the DFASTP program. An early eluting peak, M, = 28,000, produced the only unknown sequence. Further internal sequence of this protein was obtained by digesting the purified material with a 1:10 ratio (enzyme to substrate) of lysyl endopepticase (Wako Chemicals) at pH 80 in the presence of 0.05% SDS, 0.1 M Tris-HCl at 37°C for 18 hr. The digested peptices were then separated on HPLC as above. The two major eluting peaks, PF I and PF II, were sequenced as above.

cDNA Cloning

Two Agi10 libraries, a placental cDNA library prepared as described (Ullrich et al., 1985), and a random primed cDNA library made from the promyelomonocytic cell line HL-60 (provided by Karen Fisher) were probed with two oligonucleotide probes derived from the protein sequences of PFI and PFII using human codon bias (Lathe, 1985). The probes, 5'-AAGGGCACCTACCTGTACAATGACTGCCCTGGCTTTGGC-CAGGATGAGAATGA from PF I and 5'-AAGGAGATGGGCCAGGTGGA-GATCTCCICCTGCACAGTGGAC AATGACACAGTGTGTGG from PF II, were labeled with [γ-32P] ATP using 74 polynucleotide kinase and used to screen replicate sets of nitrocellulose filters containing \sim 1 x 106 independent clones from each of the two libraries. Filters were probed at 42°C in a hybridization solution that was 20% formamide, 0.1% SDS, 5x Denhard's solution, 50 µg/ml salmon sperm DNA, 50 mM NaPO4, 0.1% sodium pyrophosphate. Filters were washed twice in 0.5x SSC, 0.1% SDS at 42°C. Hybridizing phage were plaque purified, ONA was prepared, and cONA inserts were isolated and subcloned using standard techniques (Maniatis et al., 1982). Four clones from the HL-60 library, HL-60-2, -3, -10, and -14, and one clone from the placental library were sequenced on both strands using the chain termination procedure (Sanger et al., 1977).

Northern Analysis of TNF-R mRNA

Northern hybridization was performed as previously described (Thomas, 1980; Wong et al., 1988). Briefly, total cytoplasmic RNA was extracted from cells, anriched for poly(A)* mRNA, electrophoresed on a formaldehyde-agarose (1.2%) gel, and transferred to nitrocellulose. The filters were baked for 30 min at 80°C under vacuum and hybridized to a 32P-labeled TNF-R probe for 16 hr. The probe consisted of the cDNA insert isolated from the placental clone (2.1 kb EcoRl fragment) labeled with [o-32P]dATP and [o-32P]dCTP by the random priming method. Filters were washed at 60°C in 0.1× SSC, 0.1% SDS for 30 min. Autoradiography was carried out for 24 hr using Kodak XAR-5 film and an intensifying screen.

TNF-R Expression Plasmid and Transfection of TSA 201 Cells A 2.1 kb TNF-R cDNA fragment was isolated by a partial EcoRI digest

from the placental phage TNF-R clone. This fragment was ligated into the EcoRI site of the expression plasmid pRK5 (R. Klein and D. V. G., unpublished data). The cONA in the expression construct, pRK-TNF-R. is downstream of the cytomegalovirus promoter/enhancer and under its transcriptional control. Downstream of the cDNA insert are SV40 termination and polyadenylation signals. Human TSA 201 cells (obtained from R. DuBridge) are a derivative of the human embryonic kidney cell line 293s (Graham et al., 1977), which constitutively expresses large T antigen. These were transfected with either the pRK-TNF-R expression plasmid or mock transfected with the pRK plasmid without a cDNA insert. Transfections were performed in 100 mm plates using 7.5 ng of plasmid DNA per plate by the calcium phosphate precipitation method essentially as described (Gorman, 1985), except that the precipitates were left on the cells for 16-18 hr, and the cells were not shocked with DMSO in PBS. The transfected cells, transiently expressing TNF-R, were assayed 48 hr after transfection.

Analysis of Transfected TSA 201 Cells

For FACS analysis, TNF-a was biotinylated using biotin-N-hydroxysuccinimide ester at a 1:2.5 ratio of biotin ester:protein as described (Ranges et al., 1989). Forty-eight hours after transfection, cells (105) were treated with 50 nM biotinylated TNF-e by incubation for 2 hr at 4°C in PBS + 2% fetal calf serum (FCS). After washing twice the cells were stained for 30 min at 4°C with phycoerythrin (PE)-conjugated streptavidin and then washed twice and resuspended in PRS 4, 204

Cloning of a Human TNF Receptor

FCS containing 0.5 µg/ml propidium iodide. For competition binding with unlabeled TNF- α and TNF- β , cells were preincubated with either 250 nM nonbiotinylated TNF- α or 100 nM nonbiotinylated TNF- β prior to staining with PE-conjugated streptavidin plus 50 nM biotinylated TNF- α . The cells were analyzed on a Coutter Elite Flow Cytometer using the 488 nm line of an argcn ion laser, gating only on propidium iodide—excluding cells. PE emission was detected using a 575 nm (\pm 25 nm) band pass filter, and propidium iodide was detected with a 680 nm long pass filter. Electronic compensation was used to subtract the spectra overlap of PE into the propidium iodide detector.

For analysis of binding of 1251-TNF to transfected ceils, pAK-TNF-R or pRK mock-transfected cells were harvested with PBS commining 1 mM EOTA and washed with PBS containing 0.1% BSA and 0.02% sodium azide (PBSA buffer). Duplicate or triplicate samples of 2-2.5 \times 108 cells in 0.5 ml of PSSA were incubated at 4°C for 2 hr while shaking with 16.7 pM 125|-TNF-a (New England Nuclear: 88.6 mCi/mg) alone or in the presence of increasing concentrations of unlabeled recombinant human TNF-α or TNF-β (rHuTNF-α and -β, Genentech). Nonspecific binding was determined by the addition of 0.33 mM unlabeled TNF-a. Cells were centriluged at 14,000 $\, imes\,$ g for 15 min, and unbound 125 TNF a was aspirated. The cell pellet was washed once with 1 ml of ice-cold PBSA. The amount of 1251 bound was determined by counting the cell pellets in a gamma counter. The data were fit using nonlinear least-squared regression analyses according to Marquardt algorithms (GraphPAD Inplot version 30, GraphPAD Software, San Diego, CA).

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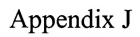
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A Tumor Necrosis Factor-binding Protein Purified to Homogeneity from Human Urine Protects Cells from Tumor Necrosis Factor Toxicity*

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Unfractionated preparations of the proteins of human urine provided protection against the in vitro cytocidal effect of tumor necrosis factor (TNF).

In certain cells, the proteins decreased expression of the receptors for TNF in a temperature-dependent way.

In all cells examined, the proteins were found to interfere also with the binding of both TNF and interleukin-1 when applied directly into the binding assays. That effect could be observed in the cold, suggesting that it was independent of cellular metabolism.

A protein which protects cells against the cytotoxicity of TNF was purified from human urine by chromatography on CM-Sepharose followed by high performance liquid chromatography on Mono Q and Mono S columns and reversed phase high performance liquid chromatography. This protein is a very minor constituent of normal urine, with an apparent molecular weight of about 27,000 in sodium dodecyl sulfatepolyacrylamide gel electrophoresis under both reducing and nonreducing conditions. Homogeneity of the purified protein was confirmed by microsequence analysis which revealed a single N-terminal sequence: Asp-Ser-Val-Cys-Pro-. The protein protected cells from TNF toxicity 2: concentrations of a few nanograms per ml and interfered with the binding of both TNF- α and TNF- β to cells, when applied simultaneously with the cytokines. However, unlike crude preparations of the urinary proteins, the purified protein did not induce in cells a decrease in ability to bind TNF nor did it interfere with the binding of interleukin-1 to its receptor. Direct, specific binding to the protein of TNF-a and, to a lesser extent, also TNF-\$\beta\$, but not of interleukin-1 nor interferon-7 could be demonstrated. It is suggested that this protein blocks the function of TNF by competing for TNF with the TNF receptor and not by interacting with the target cell.

Tumor necrosis factor (TNF)¹ is outstanding among the various mediators of immune defense in the extent to which it may cause harm to the host. Although it is effectively protective against various pathogens, this cytokine also has a mediating role in the pathological manifestations of diseases, including those caused by these very pathogens against which TNF can protect (for review, see Refs. 1 and 2). Exploring ways for suppressing the formation of TNF and antagonizing its destructive potential seems, therefore, of just as much practical importance as defining ways to take advantage of the beneficial effects of TNF in therapy.

Human urine has been shown to contain proteins which can interfere with the function of interleukin-1 (IL-1) (3-5). In view of the marked similarity in the physiological function of TNF and IL-1, we have posed the question whether proteins found in urine can also suppress the activity of TNF. We report here the purification to homogeneity and initial characterization of a protein which is present in human urine in minute amounts. This protein binds TNF, thus preventing its interaction with the TNF receptors and blocking its activity. It does not interfere with the binding of IL-1 to cells and differs in this, as well as in some other characteristics, from uromodulin, a urinary protein which suppresses the function of IL-1 (5-7) and, according to a recent study (7), also binds TNF with a high affinity, although it is apparently unable to interfere with its function. During preparation of this manuscript a study by Seckinger et al. (8) was published, describing a urinary antagonist to TNF which may be identical to the one described here.

MATERIALS AND METHODS

Cells

Murine A9 cells (9) and human foreskin fibroblasts, FS11 (established in our laboratory by Dr. D. Rotman), were cultured in Dulbecco's modified Eagle's minimal essential medium containing 100 units/ml penicillin, 100 μ g/ml streptomycin, and 2 μ g/ml amphotericin B. The media were supplemented with 10% newborn calf serum for the A9 cells and with 10% fetal calf serum for the FS11 cells.

Cytokines

Recombinant human TNF- α (rhuTNF- α , 6×10^7 units/mg protein), recombinant murine TNF- α (rmuTNF- α , 2.6×10^7 units/mg protein), and recombinant human TNF- β (rhuTNF- β , lymphotoxin, 1.2×10^6 units/mg protein) were kindly provided by Dr. G. Adolf, Boehringer Institute, Vienna, Austria. Recombinant human IL- 1α (rlL- 1α , 2.5×10^6 units/mg protein), consisting of the 154 carboxyl-

† Recipient of a fellowship from the Minerva Foundation (Federal Republic of Germany).

§ Holder of the Maurice and Edna Weiss Chair in Interferon Research.

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The abbreviations used are: TNF, tumor necrosis factor; CUP, crude urinary proteins; IFN, interferon; HPLC, high performance liquid chromatography; PAGE, polyacrylamide gel electrophoresis; TBP, TNF-binding protein; SDS, sodium dodecyl sulfate; IL-1, interleukin-1; PBS, phosphate-buffered saline; r, recombinant; hu, hu-

Appendix K

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A TNF-binding Protein from Human Urine

terminal amino acids of the 271-amino acid human IL-1 precursor, was a gift of Drs. A. Stern and P. T. Lomedico (Hoffmann La Roche, Nutley, NJ). Recombinant human interferon- γ (rIFN- γ) (5 × 10² units/mg protein) was provided by Dr. D. Novick of our laboratory.

Radiolabeling of Cytokines

All cytokines were labeled with ¹³I by the chloramine-T method, as previously described (10). The amounts of incorporated label in the preparations of cytokines used in this study were as follows: 112 µCi/µg protein for rmuTNF-a, 126 µCi/µg protein for rhuTNF-a, 113 µCi/µg protein for rhuTNF-d, and 139 µCi/µg protein for rlL-1-a. The purified TNF-binding p. tein (TBP) was labeled by the same method to a specific activity of 238 µCi/µg protein.

. Assays for the TNF-binding Protein

Quantitation of the Protect'? Effect of the TBP, against TNF Cytotoxicity—Mouse A9 cells were seeded in 96-well microtiter plates at a density of 13,000-20,000 cells/well. Urinary protein samples were applied, about 24 h later, together with cycloheximide (50 µg/ml) and rhuTNF-a (5 units/ml), and the cells were further incubated at 37 °C for 14 h. Cell viability was then quantitated by the neutral red uptake assay (11). For maximal sensitivity, the test was initiated when the A9 cells were just about to reach confluency. One unit of protective activity was defined as the amount of TNF-binding protein in whose presence the number of cells remaining viable, under the conditions of the assay, was doubled. The morphology of the A9 cells when protected from TNF toxicity by TBP is shown in Fig. 1.

Quantitation of the Binding of Cytokines to Cells and Its Decrease

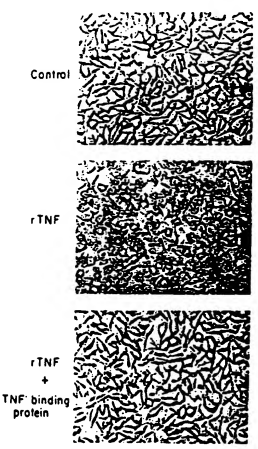


FIG. 1. The cytocidal effect of TNF in A9 cells and its decrease by the urinary proteins. Photographs of cells used for the bioassay of the protective effect of the urinary proteins (see "Materials and Methods") were taken after 14 h treatment. Top panel, control, cells treated with cycloheximide only, middle panel, dead cells after treatment with TNF + cycloheximide; bottom panel, cells treated with TNF + cycloheximide in the accesses of 250 units for the cycloheximide.

by Proteins of the Urine-A9 and FS11 cells were seeded into 15-mm tissue culture plates at a density of 2.5 × 10° cells/well. After 24 b incubation at 37 °C in 5% CO2 atmosphere, the plates were transferred to ice, the growth medium was removed, and the radiolabeled cytokines (10s cpm/plate, counting efficiency 50%) were introduced in 150 µl of ice-cold phosphate-buffered saline (PBS) (140 mm NaCl, 1.5 mm KH,PO., 8 mm Na,HPO., 2.7 mm KCl, 0.5 mm MgCl, 0.9 mM CaCl₂ containing 0.5% bovine serum albumin and 0.1% NaNa (PBS/BSA)). Following incubation for 2 h on ice, the cells were rinsed twice with ice-cold PBS/BSA and detached with Cath and Mg*-free PBS containing 5 mm EDTA. The cell-associated radioactivity was determined using a y-counter Nonspecific binding of the cytokines, determined by adding 1000-fold excess of unlabeled cytokine, was subtracted from all values. Effects of urinary proteins on the binding of cytokines were examined by either applying the proteins directly into the binding assay or pretreating the cells by the proteins, for various durations, either ... 37 °C (in growth medium) or at 4 °C (in PBS/BSA) and then removing the proteins prior to the quantitation of cytokine binding. Samples of the proteins were tested in duplicates.

Solid Phase Assay for the Binding of Various Cytokines to the Purified TNF-binding Protein—PVC 96-well radioimmunoassey plates (Dynatech 1-220-25) were coated with rhuTNF-a, rhuTNF-3, rlL-1a, or rlFN-7 by incubation, for 12 h at 37 °C, with solutions of 5 μ g/ml of the pure cytokines in PBS containing 0.02% NaN, The wells were then rinsed and incubated further, for 8 h at 4 °C, with PBS containing 0.5% BSA, 0.02% NaN, and 0.05% Tween-20 (blocking solution). Samples of radiolabeled TBP (10° cpm in 50 μ l of blocking solution) were then applied, either alone or in the presence of various cytokines or excess unlabeled TBP, and the plates were incubated for 12 h at 4 °C. They were then rinsed three times with blocking solution. The counts of the material which remained bound to the PVC plates were determined using a γ -counter.

Purification of the TNF-binding Protein

Concentration of the Crude Urinary Proteins (CUP)—Urine from healthy male donors was processed in pools of 300 liters. The urine was filtered on a Millipore HVLP membrane (pore size, 0.5 µm) using a Pellicon cassette system. The filtrate was concentrated by tangential ultrafiltration to a final volume of 750 ml, with the aid of a PTGC Millipore membrane having a molecular weight cut off at 10,000. The concentrate was dialyzed against PBS containing 0.02% NaN₃ and 1 mM benzamidine (Sigma), divided into portions, and frozen.

Chromatography on CM-Sepharose—A CM-Sepharose (Pharmacia, Uppsala, Sweden) cation exchange column (2.7 × 10 cm) was prewashed with 1 M NaCl, 10 mM citric acid, pH 5.0, containing 0.02% NaN₃ (buffer C) and equilibrated with 10 mM citric acid (pH 5) containing 0.02% NaN₃ (buffer A). The concentrate of urinary proteins was dialyzed against buffer A and centrifuged for 15 min at 8000 × g. The supernatant was applied at 4 °C on the column at a flow rate of 2 ml/min. The column was washed with 1500 ml of buffer A and eluted with 250 ml of a solution containing 200 mM NaCl, 10 mM citric acid (pH 5.0), and 0.02% NaN₃ (buffer B). A second step of elution was performed with 150 ml of buffer C. Fractions of 50 ml were collected and tested for biological activity, and their protein concentration was determined.

Cation Exchange HPLC—The active fractions eluted from the CM-Sepharose column were pooled, dialyzed against buffer A, and applied on a Mono S HR 5 × 50-mm column (Pharmacia). The column was washed at a flow rate of 0.5 ml/min until all unbound proteins were removed. The bound proteins were eluted with a linear NaCl gradient (0-350 mM) in buffer A. The gradient was run for 40 min at a flow rate of 0.5 ml/min. The column was then washed for 10 min in buffer D (350 mM NaCl in buffer A), and further with buffer C. Fractions of 0.5 ml were collected and examined for a protective effect against TNF cytotoxicity, and their protein concentration was determined.

Anion Exchange HPLC—A fast protein liquid chromatography Mono Q HR 5 × 50-mm anion exchange column (Pharmacia) was equilibrated with 5 mm sodium borate (pH 9.0) containing 0.02% NaN₃ (buffer E). The active fractions eluted from the Mono S column were pooled, dialyzed against buffer E, and loaded on the Mono Q column. The column was washed with buffer E until all unbound proteins were removed. The bound proteins were eluted at a flow rate of 0.5 ml/min, with a 30-min linear gradient from 0 to 60 mm NaCl in huffer E followed him 20 min linear gradient from 0 to 60 mm NaCl in huffer E followed him 20 min linear gradient from 0 to 60 mm NaCl in huffer E followed him 20 min linear gradient from 0 to 60 mm NaCl

buffer E. Fractions of 0.5 ml were collected and examined as above. Reversed Phase HPLC—The reversed phase HPLC column squapore RP300 (4.6 × 30 mm. Brownlee Labe), was prewashed with 0.3% aqueous trifluoroacetic acid (buffer F). The active fractions from the Mono Q column were pooled and loaded on the column. The column was washed with buffer E at a flow rate of 0.5 ml/min until all unbound material was removed; it was then eluted at a flow rate of 0.5 ml/min with a 0-20% linear gradient of acetonitrile in buffer F for 5 min followed by a 20-50% linear gradient of acetonitrile in buffer F for 5 min. The column was then washed with 80% acetonitrile in buffer F for 15 min. Fractions of 0.5 ml were collected and assayed as indicated.

SDS-Polyacrylamide Gel Electrophoresis—SDS-polyacrylamide gel electrophoresis (SDS-PAGE) on 15% acrylamide gels was performed by the method of Laemmli (12) using the Bio-Rad Minigel device (thickness of gel; 0.5 mm). Proteins in the gel were visualized by silver staining (13).

N-terminal Sequence Analysis—Samples of the purified TBP were subjected to N-terminal sequence analysis on a pulsed liquid gas phase protein microsequencer (Model 475A Applied Biosystems Inc., Foster City, CA).

Protein Determination—Protein concentrations were determined by the fluorescamine method (14), using crystalline bovine serum albumin as a standard.

RESULTS

Effects of Unfractionated Preparations of the Urinary Proteins on TNF Function—Unfractionated preparations of the proteins of human urine had a marked, concentration-dependent, protective effect against the cytotoxicity of TNF (Fig. 2A, and see also Fig. 1).

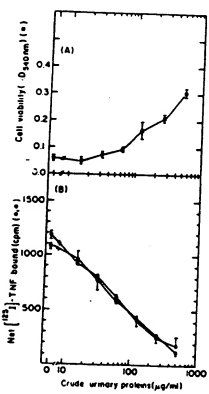


Fig. 2. Concentration dependence of the effects of unfractionated urinary protein (CUP) on TNF cytotoxicity (A) and TNF binding to cells (B). Viability of A9 cells following challenge with TNF in the presence of CUP at various concentrations (A) is demonstrated by the extent of uptake of neutral red dye by the cells (OD_{100 cm}). Effect of the proteins on the binding of ¹²³I-TNF to FS11

To explore the mechanisms which underly this protection, we examined the effect of the proteins on the binding of TNF to its receptors. This test was carried out in several different ways: (a) pretreating the cells with the urine proteins at 4 °C, prior to the quantitation of TNF binding, to find out if these proteins block the TNF receptors: (b) pretreating the cells with the proteins at 37 °C, to see if there are components in the urine which down-regulate the TNF receptors: (c) supplying the proteins, simultaneously with TNF, directly to the TNF binding assay mixture, to find out if constituents of the urine interfere with the binding of TNF by an effect on TNF itself.

Pretreatment of human foreskin fibroblast at 4 °C with the crude urinary proteins (CUP) had no effect on subsequent binding of TNF in the absence of the CUP (Fig. 3, left, top panel, 4 °C). However, after treatment with CUP at 37 °C, the cells exhibited a marked decrease in ability to bind TNF. This induced effect was rapid and transient, reaching a maximum within an hour of application of the proteins and then gradually decreasing (Fig. 3, left, top panel). A similar induced decrease in TNF binding has been reported in certain cells following treatment with IL-1 (15). Indeed, treating the human fibroblasts with IL-1, resulted in a decrease in TNF binding displaying kinetics similar to the kinetics of the effect induced by CUP (Fig. 3, left, middle panel). Murine A9 cells treated with CUP did not exhibit any such induced decrease in binding of TNF (not shown).

In addition, the CUP had a marked inhibitory effect on TNF binding when applied directly into the TNF binding

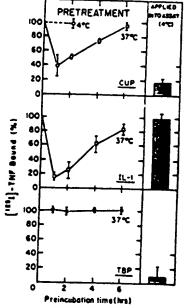


FIG. 3. Influence of the crude urinary proteins, IL-1a and pure TNF-binding protein on the binding of ¹²⁸I-TNF to FS11 cells. Left, pretreatment with the proteins; right, application of the proteins together with ¹²⁸I-TNF to the binding assay, top, effects of the crude urinary proteins (CUP) at 510 µg/ml; middle, effects of IL-1a at either 4 fg/ml (10 units/ml, left) or 4 µg/ml (right); bottom, effects of the purified TNF-binding protein (TBP) at 10 ng/ml. In the experiment presented in the left panels, cells were preincubated for the indicated duration either at 37 °C with the CUP (O), IL-1 (D), or TBP (O) or at 4 °C with CUP (O). The proteins were then removed and the binding of ¹²⁸I-TNF to the cells was quantitated as described under "Materials and Methods." In the experiment presented in the right panels, CUP, IL-1, and TBP were applied directly into the TNF-

assay (Fig. 3, right, top panel). The effect occurred also at 4 °C and could not be increased further by preincubating TNF with the CUP at 37 °C (Fig. 2B). It could be observed in all cells examined, including the human foreskin fibroblasts (Fig. 2) and murine A9 cells (Table I). The CUP also interfered with the binding of TNF-\$\beta\$ (lymphotoxin) and IL-1 to cells, when applied simultaneously with the cytokines (Table I).

Several trivial causes for the effects of urinary proteins on the activity of TNF seemed to be excluded by the above observations. The presence of TNF itself in the urine would have resulted in interference by the urinary proteins with the binding of radiolabeled TNF and, in certain cells, also in induction of resistance to TNF toxicity (16). However this possibility seemed to be excluded by the fact that the urinary proteins did not interfere with the binding of TNF when applied to cells in the cold, pric: to the application of TNF. Were TNF present in a free form in the urine, it would be expected to bind to the TNF receptors in such a pretreatment and thus block the binding of . bsequently applied radiolabeled TNF. The presence of biologically active TNF in the CUP seemed to be excluded also by the fact that these preparations did not have any toxic effect on cells, not even when applied in the presence of cycloheximide, which sensitizes cells to TNF toxicity (data not shown).

IL-1 has also been shown to induce, in certain cells, resistance to the toxicity of TNF, as well as a decrease in TNF binding (15, 17). However, unlike the urinary proteins, IL-1 does not interfere with TNF binding when applied to cells simultaneously with TNF, in the cold (Fig. 3, right, middle panel; see also Ref. 15) and, therefore, even if present in the urine, cannot account for that effect of CUP.

We also considered the possibility that the decrease in TNF binding and activity was due to degradation of TNF by some proteases known to be present in the urine. The fact that incubation of TNF with the urine proteins at 37 °C for 2 h prior to their application to the cells did not enhance the interference of the CUP with TNF binding (Fig. 2B), testifies that neither proteolytic degradation nor any other enzymic modification of TNF is involved in this effect. Furthermore, analysis by SDS-PAGE of the molecular size of TNF after its incubation with the urinary proteins revealed no signs of such degradation (not shown). Several agents known to block protease activity, N-ethylmaleimide, No-p-tosyl-L-lysine chloromethyl ketone, benzamidine, iodoacetamide, phenylmethanesulfonyl fluoride (all tested at a concentration of 1 mm), leupeptin (at 1 µg/ml), and aprotinin (at 0.5 unit/ml), did not interfere with the inhibition of TNF binding by the CUP (data not shown).

Involvement of antibodies in the effect of CUP seemed

TABLE I Interference of the urinary proteins with the binding of various cytokines to cells

The various cytokines, radiolabeled with ¹³⁸L, were applied to cells either alone, or together with the crude urinary proteins (CUP, 510 $\mu g/ml$) or the pure TNF-binding protein (TBP, 10 ng/ml). Specific binding was determined as described under "Materials and Methods." Values are given as percent of the binding of each cytokine in the absence of the inhibitory protein, which for rhuTNF- α was 2160 cpm (\pm 160); for rhuTNF- β , 1210 cpm (\pm 102); for rhuIL-1 α , 1019 cpm (\pm 691; and for rmuTNF- α , 13,543 cpm (\pm 188).

	Binding	resuTNF-a		
	rhuTNF-e	rbuTNF-8	rille	binding to A9 cells
	*	5	8	\$
Untreated cells	100	100	100	100
CUP (7 units/ml)	11 ± 5	35 ± 5	59 ± 11	22 ± 1

unlikely since, normally, urine does not contain antibodies. Furthermore, the molecular size of the proteins which mediate the effect on TNF activity, as estimated by size exclusion chromatography (see below), is clearly lower than that of immunoglobulins.

Purification and Initial Characterization of the Urinary TNF-binding Protein—A bioassay for the urinary protein(s) which mediate a protective effect against TNF cytotoxicity was established (see "Materials and Methods"); it then was used for the detection of these protein(s) throughout various fractionation steps.

Several chromatographic approaches for purification of the protein(s) were attempted. In size exclusion chromatography (on an Ultrogel AcA44 column equilibrated with phosphatebuffered saline) only poor resolution between the protein(s) which interfere with TNF activity and other urinary proteins could be observed. Under these conditions, the activity fractionated together with the majority of the protein mass, peaking at an apparent molecular size of about 50,000-70,000. On the other hand, in isoelectric focusing, some enrichment of the proteins could be obtained. Consistent with prior observations, much of the protein mass of urine was found to be rather acidic, with an apparent isoelectric point lower than 5.0. Yet the isoelectric point of the protein(s) which protect cells from TNF toxicity was found to be close to 6.0 (data not shown). Fractionation by isoelectrofocusing for initial enrichment of the protein seemed impractical since only limited amounts of protein can be applied at a time. However, the pattern of isoelectric points revealed by that procedure indicated that enrichment of the protective protein on the basis of its charge properties should be possible.

As a first step, the urinary proteins were fractionated at pH 5.0 on a carboxymethyl-Sepharose column. Consistent with their acidity, most of the proteins did not bind to the resin under those conditions. However, the inhibitor of TNF activity was bound effectively to the resin at that step (about 80% of the applied activity), and most of it could be eluted from the column, together with about 1% of the initially applied protein, by increasing the ionic strength by 0.2 M NaCl. Increasing the ionic strength further (1 M NaCl) did not result in elution of any additional activity.

In the second purification step, the proteins were fractionated on a cation exchange HPLC column. The active protein was eluted at about 180-220 mm NaCl (Fig. 4). Peak fractions

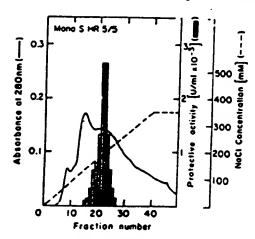


FIG. 4. Mono S cation exchange HPLC of TBP. The column was equilibrated with 10 mM citric acid (pH 5), 0.02% NaN₂. Urinary proteins, enriched for TBP activity by fractionation on CM-Sepharose, were applied and then eluted with NaCl gradient (---). The

were pooled and subjected to further purification on an anion exchange HPLC column, from which the TNF inhibitory activity was eluted at a salt concentration of about 40 mM (Fig. 5).

The final fractionation step was on an Aquapore RP 300 reversed phase HPLC column. Proteins were eluted from the column by applying a gradient of acetonitrile. The active protein was found to elute as a distinct protein peak, at about 27% acetonitrile (Fig. 6).

Analysis by SDS-PAGE, revealed that the factor was purified at that step to homogeneity; it was identified as a protein with an apparent molecular weight of about 27,000 (inset in Fig. 6). The molecular size was independent of whether or not the analysis was performed in the presence of a reducing agent (ρ -mercaptoethanol). Homogeneity of the purified protein was further confirmed by N-terminal microsequence analysis. A single sequence, Asp-Ser-Val-Cys-Pro-, was obtained in the analysis at a high yield (initial yield, 67%).

Both from the low amounts of the protein, recovered in the purification, and from comparison with the protein pattern in the crude preparation, as revealed by SDS-PAGE, it is clear that the factor is a very minor constituent of urine. In the urine preparation, used for the fractionation whose results are presented in Table II, the protein constituted 10⁻³~10⁻⁴% of the total protein. Specific activity of the purified protein was about 50,000-fold higher than that of the CUP.

Like the crude preparations of the urinary proteins, the purified protein interfered with the binding of TNF- α (of both human and mouse origin), as well as with huTNF- β when applied to cells simultaneously with these cytokines (Table I and Fig. 3, right, bottom panel). However it did not interfere, to any measurable extent, with the binding of IL-1 to its receptor (Table I) nor did it induce in cells, pretreated with the protein, a decrease in ability, to bind TNF (Fig. 3, left, bottom panel). The latter two effects of crude preparations of the urinary proteins therefore seem to be mediated by some other constituents of the urine.

The binding properties of the urinary protein were explored using radiolabeled preparations of the purified protein. As shown in Table III, the labeled protein was found to bind to immobilized TNF- α and, to a much lesser extent, to TNF- β . This binding could be competed by TNF- α and TNF- β and by excess of the unlabeled urinary protein. IL-1 (as well as

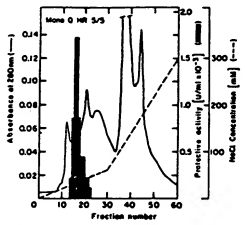


FIG. 5. Mono Q anion exchange HPLC of TBP. Active fractions eluting from the Mono S column were made up to 5 mM sodium

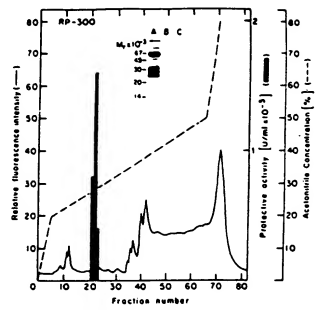


Fig. 6. Reversed phase HPLC of TBP. The proteins enriched for TBP activity on the Mono Q column were applied to an Aquapore RP300 column. Elution was performed with a gradient of acetonitrile in 0.3% aqueous trifluoroacetic acid (- - -). Fractions were examined for bioactivity (2), and protein (-—) content. Shown in inset is SDS-PAGE analysis of proteins in the absence of reducing agents (A. B), or in the presence of 15% β -mercaptoethanol (C). The pettern of proteins in the preparation of unfractionated urinary proteins prior to chromatography on CM-Sepharose (5 µg. lone A), is compared to the protein in fraction 21 of the elution from the HPLC RP300 column (0.1 µg, lanes B and C), where only a single polypeptide band can be discerned. (The faint high molecular weight bands in C could be observed also in the absence of any added protein and seem to reflect the presence of some contaminants in the θ -mercaptoethanol) By comparing to the migration of molecular weight markers (Pharmacia, shown in the left), the molecular weight of the purified TBP was estimated as about 27,000.

TABLE II
Purification of the TNF-binding protein

Purification step		Protective activity		Specific	Puri	Purification	
	Protein			activity	Per	Total	
	TV.	white	%	units/mg		fold	
Starting material (CUP)	7.1 × 10°	90,100	100	13			
Carboxymethyl- Sephaross	8.9 × 10 ¹	54,400	60	610	48	48	
Mono S	1.7×10^{1}	20,000	22	1,120	1.8	88	
Mono Q	4.8 × 10 ⁻¹	13,000	14	29,100	24.3	2.100	
HPLC RP 300	1.1 × 10 ⁻²	7,000	8	600,000	23.8	54,000	

IFN- γ) did not bind the urinary protein, nor did it compete for the binding of the protein to TNF.

DISCUSSION

A protein present in the human urine in minute amounts is shown in this study to interfere with the function of TNF by blocking the binding of TNF to its receptors. In the crude state, the urinary proteins suppress the binding of TNF to cells by effects on both the cell and the TNF molecules; they also interfere with the interaction of cells with IL-1-cytokines

TABLE III

Binding of the TNF-binding protein to cytokines and the effect of competitive proteins

Binding of purified, radiolabeled TBP to the cytokines listed in the left column and cross-competition with the cytokines indicated or with excess unlabeled TBP (last column) was determined in a solid phase assay as described under "Materials and Methods." The concentrations of the proteins applied for competition for the binding to TBP were as follows: TNF-a, 5 µg/ml; TNF-3, 1 µg/ml; IL-1a, 4 µg/ml; IFN-7, 1 µg/ml; TBP, 80 µg/ml.

Cytokine examined for			Proteins ap	plied for competition for	TBP binding	
bind	ling to TBP	TNF-a	TNF-3	IL-la	IFN-7	ТВР
TNFa TNFJ IL-la IFNy	17,820 (±130) 450 (±40) 140 (±40) 110 (±10)	400 (±10) 130 (±10) 130 (±30) 100 (±10)	14,340 (±900) 240 (±30) 105 (±10) 105 (±10)	cpm (**//TBP bound 18,520 (±450) 480 (±30) 100 (±10) 90 (±10)	17,810 (±700) 440 (±20) 130 (±30) 120 (±20)	790 (±20 110 (±%) 130 (±30 120 (±10

crude urine against TNF toxicity, seems to function only by affecting the TNF molecules; it showed no direct effect on cells and was unable to interfere with the binding of IL-1 to its receptors.

The purified protein acts by binding the cytokine and thus competing for it with the TNF receptors. Although it does not bind IL-1, it does bind TNF- α and, with much lower effectivity, also TNF- β (lymphotoxin). The specific nature of its interaction is further demonstrated by the inability of the protein to bind another cytokine-IFN- γ . Even though TNF- α and TNF- β share only partial structural homology (18), they compete for binding to the same cell surface receptor (19). The fact that they both bind to the urinary protein raises the possibility that this protein associates with that part of the cytokine molecule which is recognized by the receptor.

By its inability to interfere with the binding of IL-1 to cells, the TNF-binding protein (TBP) can be distinguished from uromodulin, a major glycoprotein of the urine, of greater molecular size (85,000), which was reported to have a high binding affinity to both IL-1 and TNF and to interfere with the function of IL-1, although not with that of TNF (5-7). The TBP is also clearly distinct from another antagonist to IL-1, shown recently to be present in urine, which interferes with IL-1 binding apparently by binding competitively to the IL-1 receptor, but it seems unable to block the function of TNF (4, 20, 21).

The presence of the above antagonists in urine probably accounts for the inhibitory effect of CUP on IL-1 binding. At the same time, the induced decrease in the ability of FS11 cells to bind TNF following treatment with CUP appears to be very similar to an effect of IL-1 itself. IL-1 induces a decrease in binding of TNF, which apparently reflects a decrease in expression of the TNF receptors. In FS11 fibroblasts, although not in some other cells (15), this decrease is transient and its kinetics resembles the one seen with urinary proteins. Some evidence for the presence of IL-1 in the urine has been reported (4, 22). Whether indeed IL-1, if present in the unfractionated preparations of the urine, can mediate the effect in spite of its coexistence with proteins which block its activity, or whether it is some other constituent(s) of the urine which functions in our test system similarly to IL-1, remains to be determined.

The protein whose purification is described in this study is present in the urine in minute amounts, constituting about $10^{-3}-10^{-4}\%$ of the total protein mass. It would not have been detected save for the fact that it is very active. Indeed, it can be calculated that a unit of protective activity (the activity resulting in a 2-fold increase in the number of cells which remain viable after challenge with TNF) is mediated by the

The physiological role of the protein remains to be elucidated. It is tempting to speculate that, just as in our in vitro experimental system, this protein actions vivo as an antagonist to TNF. However, to test this hypothesis it will be necessary first to determine how this protein is formed and what results from its interaction with TNF under in vivo conditions. At present, one cannot rule out that this protein has a completely different in vivo role, perhaps even a converse one; it might prolong the exposure of the organism to endogenously produced TNF by binding the cytokine and then releasing it slowly in an active form.

There is some evidence both for the existence of mechanisms whereby the organism can protect itself from the potentially destructive effects of TNF and for enhanced expression of these mechanisms following exposure to TNF. Several studies have shown that vulnerability to certain deleterious effects of TNF and of bacterial components which can induce TNF production is markedly reduced following exposure of the organism to TNF itself or to IL-1 (23-26). If the protein described in the present study can indeed contribute to such protection, it is very likely that useful applications for it will be found, specifically as a therapeutic agent in those pathological situations where TNF can have detrimental effects.

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Addendum—Recently, we have determined the sequence of the 18 N-terminal amino acids of the TBP. A search through the NBRF protein data bank, release 18, failed to reveal significant homology to any of the other known protein sequences. In the 2nd International Conference on TNF and Related Cytokines held January 15-20, this year, I. Olsson from the University of Lund, Sweden, also reported on the purification of a urinary protein which binds TNF. Based on comparison of N-terminal amino acid sequences and of chromatographic properties, this protein appears identical to the one whose purification is described in the present study.

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Appendix L

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Mechanisms Which Take Part in Regulation of the Response to Tumor Necrosis Factor

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In view of increasing evidence for a causative role of TNF in the pathogenesis of certain diseases (1,2), there is particular interest in elucidation of the mechanisms by which the function of this cytokine is controlled. The studies described below provide initial information on two different reflections of those mechanisms. One, concerns a mechanism which may contribute to a marked sensitization to some deleterious effects of TNF, observed in animals which were afflicted with pathogens (3-6). The other was initiated in an attempt to shed light on the mechanisms by which a decrease in responsiveness to TNF, such as observed following preexposure to a low dose of TNF, may occur (7,8).

Pathogen-mediated Enhancement of the Prostaglandin Inducing Effect of TNF

Vulnerability of experimental animals to the lethal effect of injected TNF was found, in a number of studies, to increase by certain pathogenic agents, like bacteria, the malaria parasites and tumors (3-6). Other studies (e.g. cf. 9) suggested that some deleterious effects of TNF are precipitated by prostaglandins, produced in response to it. A possible mechanistic link between the above observations was indicated in a study in which we examined the effect of TNF on cells after their infection with Chlamydia trachomatis. As shown in Table 1 growth of the Chlamydia in HEp-2 cells is significantly inhibited by rTNF (see also 10). An even further inhibition is observed when treating the cells with TNF together with IFN-1, at concentrations of IFN at which IFN alone has only little effect (10,11). It has been reported that the growth of chlamydiae is inhibited also by prostaglandins (12). We therefore examined the production of prostaglandins in infected cells which were treated with TNF. Similarly to various other cells (13,14) uninfected HEp-2 cells respond to TNF in some production of PGE2 though to only a little extent. Infection with Chlamydia, which by itself results in only little induction of PGE, potentiates the TNF effect (Table 1). Maximal induction of PGE, is observed at concentrations of TNF which effectively inhibit the chlamydial growth. Treating the cells with TNF and IFN-7 together, which practically abolishes the growth of chlamydia results in a similar, pronounced induction of PGE,

TABLE 1:

Effect of TNF-x and Chlamydia on the Production of PGE2 in the HEF-2 Cells

	Chlamydial yield	PGE ₂
•	(IFU/ml)	(pg/ml)
Control	-	54
Chlamydia	$(5.1\pm0.8) \times 10^6$	510
TNF	. -	405
Chlamydia + TNF	220 ± 92	42000

Where indicated, cells were infected with chlamydia trachomatis (L2/434/Bu) at a multiplicity of infections of 1 and, following adsorption of the bacteria, treated with rINF- α at 500 ng/ml. Chlamydial yield and the amounts of PGE2 in the growth medium were determined at 48 h after infection.

Such a potentiative effect of a pathogen of an effect of TNF which can be inhibitory to its growth, provides an interesting demonstration for a mechanism by which a protective function of TNF can be adjusted to the need. Yet, when occurring in excess, the augmented production of PCE2 at the time of exposure to TNF may just as well contribute to the precipitation of the deleterious effects of this cytokine.

Isolation from the Human Unine of a TNF Binding Protein with an Inhibitory Effect on TNF Activity

Trying to identify regulatory molecules which may contribute to the decrease in responsiveness to TNF (7,8) we followed prior studies which indicated that the function of IL-1 can be suppressed by certain secreted inhibitors. Human urine has been shown to contain such inhibitors of IL-1 activity (15-17). We therefore examined whether the urine contains also components which can affect cell response to TNF. As demonstrated in Fig. 1, concentrated preparations of the urinary proteins indeed suppressed effectively the cytocidal activity of TNF (18,19). A similar observation has been reported also by others (20,21).

Examining further the effect of these proteins on the binding of radiolabelled TNF to cells, we found that in certain cells unfractionated preparations of the proteins of human urine induce, in a temperature dependent way, a rapid, though transient decrease in the expression of the TNF receptors; quite similarly to an effect which IL-1 can have in those cells (23 and see also Fig. 2). The proteins of urine did not interfere with the binding of TNF when applied to cells in the cold and then removed, prior to the application of radiolabelled TNF, ruling out that their effect reflects the function of a protein which binds to the TNF receptors competitively with TNF. However, distinctly from the way IL-1 can affect TNF binding, the urinary proteins did simultaneously with it (Fig. 2).

Applying a series of chromatographic steps we isolated from the urine the

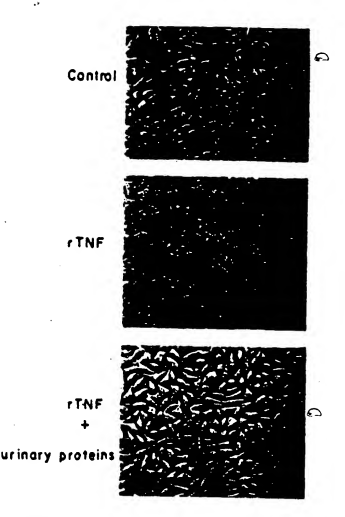
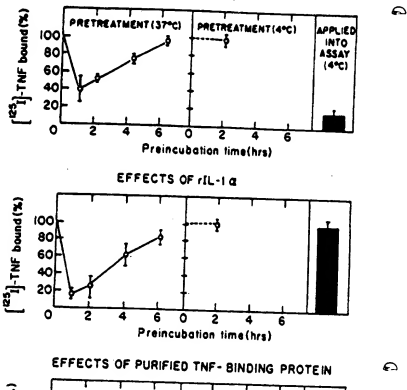


Figure 1. Protective effect of the human urinary proteins against the cytocidal effect of TNF. Top panel: Control. A9 cells treated for 14h with cyclohexamide only (CHI, 50µg/ml). Middle panel: dead cells after treatment with rTNF-cx (5U/ml) + CHI. Bottom panel: cells treated with TNF+CHI in the presence of a preparation of the urinary proteins.

protein which exarts the latter effect. It turned to be a very minor constituent of the normal urine (about 10^{-3}) of the proteins). Unlike the crude preparations, the purified protein did not induce a decrease in the expression of TNF receptors in tissue cultured cells. It was found to affect TNF by binding to it. It also binds TNF- β (LT), though at lower effectivity but not IL-1 nor IFN- γ . The protein, whose apparent molecular weight in SDS PAGE analysis was found about 30 KDa, appears, by its amino acid sequence, to be distinct from all previously isolated proteins (19).

The cellular source of this urine-derived TNF-binding protein remains to be elucidated. A likely possibility, in view of the specific nature of its binding activity, is that this protein is a shed fragment of the cell surface TNF receptor. Indeed the urine has been shown to contain a soluble form of the IL-2 receptor, which in potential may impede the function of this cytokine (23). It is tempting to speculate that the formation of the soluble TNF binding protein constitutes a way of restraining the function of TNF. Whether this indeed is its physiological role or not, the possible use of the protein in therapy for blocking the function of TNF, at times when it contributes to pathogenicity, seems worthy of exploring.

EFFECTS OF CRUDE URINARY PROTEINS ON THE BINDING OF THE TO FSII CELLS



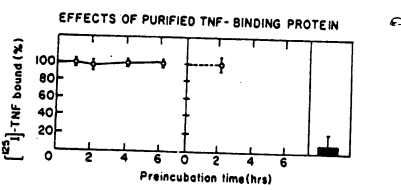


Figure 2. Influence of the grude urinary proteins, IL-1x and pure TNF binding protein on the binding of 1251-TNF to FS11 cells. Left and middle panels - Pretreatment with the proteins at 37°C and 4°C respectively. Right - Application of the proteins together with 1251-TNF to the binding assay. Top: Effects of the crude urinary proteins (CUP) at 510 µg/ml. Middle: Effects of IL-1a at either 4 pg/ml (10 u/ml-left and middle) or 4 ug/ml (right). Bottom: Effects of the purified TNF binding protein (TBP) at 10 ng/ml. In the experiments presented in the left and middle panels cells were preincubated for the indicated duration either at 37°C or at 4°C with the CUP, IL-1 or TEP. The proteins were then removed and the binding of 1251 TNF to the cells was quantitated. In the experiment presented in the right panels CUP, IL-1 and TEP were applied directly into the TNF-binding assay mixture (assay performed at 4°C). Specific binding of TNF in the absence of CUP, TEP or IL-1 (100%) was

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Comparison of sequences provided in D18.1 (Fig. 4) and D18.2 (Fig. 1D)

Jeletion or substitution in nucleic acid sequenceT: Deletion or substitution in amino acid sequence

ATC ATC Tyr Tat TAT Tyr Lys AAA AAA Lys GGA Gly GGA Gly Gln Gln CCC CAA CCC CAA Pro (Pro TCT Cys TCT GTG GTG Val Val Glu Lys Arg Asp Ser GAG AAG AGA GAT AGT GAG AAG AGA GAT AGT Ser Glu Lys Arg Asp Asp Arg GAC AGG GAC AGG Asp Arg Gly GGG Gly 999 CTA CIA Leu Pro His Leu His CAC CAC Val Pro I GTC CCT G Val Pro ↓ ↓↓↓ • Gly Leu Va • CTG r Pro ↑ ပ္ပ Ile ATT ATT Ile ↓↓↓ ↓↓↓
G1y Va1
GGG GTT
TAT CGA
TYF AFG

↑

GGC CCG GGC CCG Gly Pro Pro Cys Pro CCA TGT CCA Cys Asn Asp (Asn Asp AAT GAC Tyr TAC TAC Leu Tyr Leu TTG TTG TTC Phe ↑ Tyr \mathbf{Thr} ACC ACC Thr GGA CCA Gly Gly His Lys CAC AAA His Lys His AAG TGC (Lys Cys I Lys Cys | AAG TGC | ACC Thr ATT TGC TGT ACC Cys Thr Cys TGT Cys TCG ATT TGC cys Ile TCG Ser Asn AAT AAT Asn Asn AAT AAT Asn Gln CAA CAA Gln Pro CCT CC Pro CAC S

Thr Leu Gln Lys Pro Pro Gln Thr Leu Pro Gln Phe Thr Ala Ser Glu Asn His Leu Arg His Cys Leu TTC ACC GCT TCA GAA AAC CAC CTC AGA CAC TGC CTC TCC TTC AC- GCT TCA G-A AAC CAC CTC AGA CAC TGC CTC Phe Ser TCC Ser Ser Gly AGC GGC 9 AGC Ser Arg Glu Cys Glu AGG GAG TGT GAG GAG Glu TGT Cys AGG GAG Cys TGC cys Thr Asp ACG GAC ACG GAC Thr Asp Asp Asp GAT GAT Gln Asp CAG CAG 61y 666 666 61y

Leu Leu Ala Gln Trp Thr Gly Thr Pro Cys Val Gln Val Glu Ile Ser Ser Cys Thr Val Asp Arg Asp Thr Val Cys CAG GTG GAG ATC TCT TGC ACA GTG GAC CGG GAC ACC GTG TGT A-C TCT TCT TGC ACA GTG GAC CGG GAC ACC GTG TGT Thr ↑; CAG -TG -AG glu Ser Ser Cys Ser Lys Cys Arg Lys Glu Met Gly
AGC TGC TCC AAA TGC CGA AAG GAA ATG GGT
AGC TGC TCC AAA TGC CGA AAG GAA -TG G-T
Leu Leu Gln Met Pro Lys Gly Met Val

↑ ↑ ↑ ↑ ↑ ↑ ↑ ↑ Gly Cys Arg Lys Asn Gln Tyr Arg His Tyr Trp Ser Glu Asn Leu Phe Gln Cys Phe Asn Cys Ser Leu Cys Leu GGC TGC AGG AAC AAC CAG TAC CGG CAT TAT TGG AGT GAA AAC CTT TTC CAG TGC TTC AAT TGC AGC CTC TGC CTC GGC TGC AGG AAG AAC CAG TAC CGG CAT TAT TGG AGT GAA AAC CTT TTC CAG TGC TTC AAT TGC AGC CTC TGC CTC Ala Ala Gly Arg Thr Ser Thr Gly Ile Ile Gly Val Lys Thr Phe Ser Ser Ala Ser Ile Ala Ala Ser Ala Se

Asn Gly Thr Val His Leu Ser Cys Gln Glu Lys Gln Asn Thr Val Cys Thr Cys His Ala Gly Phe Phe Leu Arg AAT GGG ACC GTG CAC CTC TCC TGC CAG GAG AAA CAG AAC ACC GTG TGC ACC TGC CAT GCA GGT TTC TTT CTA AGA AAT GGG ACC GTG CAC CTC TCC TGC CAG GAG AAA CAG AAC ACC GTG TGC ACC TGC CAT GC- G-A TTC TTT CTA AGA Mat Glv Pro Cvs Thr Ser Pro Ala Arg Asn Arg Thr Pro Cys Ala Pro Ala Met Arg Phe Phe Leu Arg Met Gly Pro Cys Thr Ser Pro Ala Arg Arg Asn Arg Thr Pro Cys Ala Pro Ala Met Arg

GAG GAG Glu Glu Cys Val Ser Cys Ser Asn Cys Lys Lys Ser Leu Glu Cys Thr Lys Leu Cys Leu Pro Gln Ile GAG TGT GTC TCC TGT AGT AAG AAA AGC CTG GAG TGC ACG AAG TTG TGC CTA CCC CAG ATT GAG TGT TCC TGT AGT AAC TGT AAA AGC CTG GAG TGC ACG AAG TTC TGC CTA CCC CAG ATT GLU Cys Val Ser Cys Ser Asn Cys Lys Lys Ser Leu Glu Cys Thr Lys Leu Cys Leu Pro Gln Ile Glu Asn GAA AAC GAA AAC G

Leu TTA Leu CTT Leu CIT Cys TGC TGC Cys Leu CTT Phe Gly I TTT GGT C TTT GGT C Phe TTC TTC Phe thr Thr Val Leu Leu Pro Leu Val Ile Phoc CC ACA GTG CTG TTG CCC CTG GTC ATT TTC CC ACA GTC GTG TTG AAA ATG GTC ATT TTC IT Thr Val Val Leu Lys Met Val Ile Phe Gly Thr 3 GGC ACC 1 GGC ACC 2 Gly Thr 3 Glu Asp Ser G GAG GAC TCA G GAG GAC TCA G Glu Asp Ser G Val Lys Gly Thr C GTT AAG GGC ACT C GTT AAC GGC ACT C Val Lys Gly Thr C AAT AAT

Ser Leu Leu Phe Ile Gly Leu Net Tyr Arg Tyr Gln Arg Trp Lys TCC CTC CTC TTC ATT GGT TTA ATG TAT CGC TAC CAA CGG TGG AAG TCC CTC CTC TTC ATT GGT TTA ATG TAT CGC TAC CAA CGG TGG AAG Ser Leu Leu Phe Ile Gly Leu Met Tyr Arg Tyr Gln Arg Trp Lys

Appendix N

Three human transforming genes are related to the viral ras oncogenes

(human tumor cells/molecular cloning/gene families)

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Three distinct transforming genes present in human tumor cell lines are all related to the viral oncogenes of Harvey and Kirsten murine sarcoma viruses, designated v-H-ras and v-Kras, respectively. The transforming gene of a bladder carcinoma cell line has been shown to be a human homolog to v-H-ras [Parada, L. F., Tabin, C. J., Shih, C. & Weinberg, R. A. (1982) Nature (London) 297, 474-478; Santos, E., Tronick, S. R., Aaronson, S. A., Pulciani, S. & Barbacid, M. (1982) Nature (London) 298, 343-347]. The transforming gene common to one colon (SK-CO-1) and two lung carcinoma (SK-LU-1 and Calu-1) cell lines is the same human homolog of v-K-ras as is the transforming gene previously identified in a lung carcinoma cell line Lx-1 [Der, C. J., Krontiris. T. G. & Cooper, G. M. (1982) Proc. Natl. Acad. Sci. USA 79, 3637-3640]. The transforming gene of SK-N-SH neuroblastoma cells is weakly homologous to both v-H-ras and v-K-ras. NIH 3T3 cells transformed with the SK-N-SH transforming gene contain increased levels of a protein serologically and structurally related to the protein products of the v-H-ras and v-K-ras genes. Therefore, it represents a third member of the ras gene family, which we have called N-ras. Based on the homology with the v-ras genes. we have established the orientation of transcription and approximate coding regions of the cloned human K-ras and N-ras genes.

The progression of a cell from normalcy to malignancy may be due in part to the activation of transforming genes of cellular origin. The existence of cellular transforming genes has been demonstrated by the ability of genomic DNAs from certain tumors and cell lines to induce foci of transformed NIH 3T3 cells after DNA-mediated gene transfer. Transforming genes in rodent (1, 2) and human (3–9) tumor cells have been detected in this way. We have detected three distinct transforming genes in our study of 21 human tumor cell lines: one common to two lung carcinoma (SK-LU-1 and Calu-1) and colon carcinoma (SK-CO-1) cell lines, one in a bladder carcinoma (T24), and one in a neuroblastoma (SK-N-SH) cell line (9).

Several research groups have shown that certain transforming genes detected by transfer to NIH 3T3 cells are related to viral oncogenes. Der et al. (5), Parada et al. (10), and Santos et al. (11) have demonstrated that the transforming gene of T24 and EJ, two human bladder carcinoma cell lines that probably are derived from the same source (unpublished data), is the human homolog of v-H-ras, the oncogene of the Harvey sarcoma virus. Der et al. (5) have also shown that the transforming gene of Lx-1, a human lung carcinoma cell line, is a human homolog of v-K-ras, the oncogene of the Kirsten sarcoma virus. The genes that we have isolated from human tumor cell lines are related also to the viral oncogenes, designated v-onc. We demonstrate

that the transforming gene common to Calu-1. SK-LU-1. and SK-CO-1, like the transforming gene in Lx-1 characterized by Der et al. (5), is a human homolog to v-K-ras. We also demonstrate that the transforming gene of SK-N-SH is related to both v-K-ras and v-H-ras and probably codes for an immunologically crossreactive and structurally related protein. Based on the homology with the v-ras genes, we have established the orientation of transcription and probable coding regions of these genes.

MATERIALS AND METHODS

Human Tissue and Tissue Culture Cell Lines. T24. Calu-1 SK-LU-1. SK-CO-1. and SK-N-SH are human tumor cell lines (9). HT14B is a NIH 3T3 cell line transformed by Harvey sarcoma virus unintegrated viral DNA. Other transformed cell lines are described in the text.

Preparation of DNA. DNA was prepared from tissue culture cells by NaDodSO₄/proteinase-K lysis and phenol/chloroform extraction as described (9). Plasmid and bacteriophage DNAs were prepared as described (12, 13).

Enzymes. Restriction endonucleases were purchased from New England BioLabs and Bethesda Research Laboratories and used according to suppliers' instructions. E. coli DNA polymerase I was purchased from Bethesda Research Laboratories, and pancreatic DNase I was from Worthington Biochemicals.

Southern Filter DNA Blot Hybridization. DNA samples were digested with restriction endonucleases and subjected to agarose gel electrophoresis and filter-blot transfer by the method of Southern (14). Filter-blotted DNAs were hybridized with a nicktranslated 32P-labeled DNA probe under two sets of conditions. Stringent hybridization conditions entailed hybridization in a mixture containing 6× NaCl/Cit (1× NaCl/Cit is 0.15 M NaCl/ 0.015 M Na citrate, pH 7.0), Denhardt's solution (0.02% polyvinylpyrrolidone/0.02% Ficoll/0.2% bovine serum albumin . and denatured salmon sperm DNA (20 µg/ml) for 16 hr at 74°C (15), followed by sequential washing at 74°C with 2×, 1×, and 0.5× NaCl/Cit in 0.1% NaDodSO4. Nonstringent hybridization conditions entailed hybridization in a mixture containing 30% (vol/vol) formamide, 6× NaCl/Cit, 2× Denhardt's solution, E. coli DNA (100 μ g/ml), yeast RNA (200 μ g/ml), 50 mM sodium phosphate (pH 7), and 10 mM EDTA at 37°C for 36 hr, followed by washing at 50°C in 6× NaCl/Cit/0.1% NaDodSO4. Hybridized DNA was revealed by autoradiography.

Immunoprecipitation of Cellular Protein with Rat Anti-ras p21 Antiserum. NIH 3T3 normal and transformed cells were

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Abbreviations: NaCl/Cit, 0.15 M sodium chloride/0.015 M sodium citrate, pH 7.0; kbp, kilobase pairs.

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labeled in methionine-free medium containing 20 μ Ci (1 Ci = 3.7 × 10¹⁰ Bq) of [35S]methionine per ml (New England Nuclear) for 18 hr. Labeled cells were lysed in phosphate-buffered saline containing 1% Triton X-100, 0.5% sodium deoxycholate, 1 mM phenylmethylsulfonyl fluoride, and 2 units of aprotinin per ml, and the lysates were sheared through a syringe and clarified at 100,000 × g for 45 min at 4°C. Clarified supernatants were preabsorbed with goat anti-rat IgG and Staphylococcus aureus protein A. Immunoprecipitation was performed with antiv-H-ras p21 rat monoclonal antibody Y13-259 (16) (the gift of M. Furth and E. M. Scolnick) for 5 hr at 4°C, followed by addition of goat anti-rat IgC for 1 hr. Immune complexes were absorbed to protein A. and the protein A suspension was washed extensively in lysis buffer. Protein A pellets were boiled in Na-DodSO4 sample buffer and analyzed by NaDodSO4/polyacrylamide gel electrophoresis by the method of Blattler et al. (17). Radiolabeled proteins in gels were visualized by fluorography.

Molecular Clones. Clones of avian and mammalian v-onc genes are described in Table 1. λT24 and λP3 are clones from λL47.1 genomic libraries containing the transforming gene of the T24 cell line and the nontransforming homologous human sequences, respectively (25), pT24 is a pBR322 derivative with a 6.2-kilobase-pair (kbp) BamH1 insert bearing the T24 transforming gene. The SK-N-SH transforming gene is contained on a Charon 4A recombinant phage, λNPS-1-1-1 as described (26), (See also Fig. 4B.) λNPS-1-1-1 also contains portions of pBR322 and the E. coli tRNA sup F gene.

Cloning the Transforming Gene of Calu-1. An initial DNA clone of part of the transforming sequences of the Calu-1 transforming gene was obtained by using the strategy of Gusella *et al.* (27) as described by others (28, 29). DNA was prepared from NIH 3T3 secondary and tertiary transformants containing the

Table 1. Molecular clones of v-onc genes

v-onc desig- nation	Virus of origin's	Molecular clone	Restriction fragments bearing v-onc	Ref. no.
fps [‡]	PRCII SV	pRCII-1B	Kpn 1 1.5 kbp	M. Bishop [‡]
ves [†]	Y73 SV	λΥ73-11A	Sst I 4.0 kbp	18
rel [†]	ARVT	prel	EcoRI 0.8 kbp	19
ski [†]	SKV	pvski-1	Xho 1 2.8 kbp	Unpublished data
abl [§] .	Abelson MuLV	pABsub3	HindIII/Sst II 2.0 kbp	20
fes§	Feline SV	pGA FeSV	Pst I 0.5, 0.55 kbp	21
mos§	Moloney MuSV	pmos-1	Pst I 0.45 kbp	D. Dina [±]
H-ras§	Harvey	pBS-9	EcoRI 0.5 kbp	22
	MuSV	pHB-11	EcoRI/BamHI 2.2 kbp	22
K-ras§	Kirsten	pHiHi-3	EcoRI 1.0 kbp	22
11-103-	MuSV	pKBE-2	EcoRI/BamHI 3.1 kbp	22
fms§	McDonough feline SV	λSM- FeSV	<i>Kpn</i> I 2.8, 4.8 kbp	23
sis§	Simian SV	pvsis	EcoRI/Sal I 2.1 kbp	24

The table lists the v-onc sequences that were tested for homology to three human transforming genes.

transforming gene of Calu-1. Phage libraries were prepared from these DNAs in λ Charon 4A (30) by the method of Hohn and Murray (31) and screened for the presence of human sequence by the method of Benton and Davis (32); the probe was "BLUR8", a clone of the dispersed, repeated human "Alu" family sequences (33). One λ phage clone, λ L2-34, was isolated this way. Unique sequence DNAs from this clone were then used as probes for isolating "contiguous" DNA from our λ Charon 4A libraries. More than 20 independent phages containing inserts with overlapping restriction endonuclease maps were isolated in this manner. A representative set of five overlapping phage isolates, together with a composite restriction endonuclease map of 26 kbp of cloned DNA, is shown in Fig. 4A. pLC3 is the 3.0-kbp EcoRI fragment of λ L2-11 cloned into the EcoRI site of pBR322 (see Fig. 4A).

RESULTS

Three Human Transforming Genes Have Homology to Viral ras Genes. Molecular clones of v-onc were cleaved with restriction endonucleases to separate v-onc and vector sequences. and triplicate aliquots of these digests were subjected to agarose gel electrophoresis and Southern nitrocellulose filter blotting. The three replica filters were hybridized at low stringency to ³²P-labeled recombinant DNAs containing all or part of the three different human transforming genes (Fig. 1). The transforming human genes were those isolated from the bladder carcinoma cell line T24 (Fig. 1B), the lung carcinoma cell line Calu-1 (Fig. 1C), and the neuroblastoma cell line SK-N-SH (Fig. 1D).

All three human transforming genes showed homology to v-H-ras and v-K-ras (Fig. 1, lanes 8–11). The human transforming genes were not homologous to nine other v-onc genes (Fig. 1, lanes 1–7, 12, and 13). The hybridization detected in other lanes of this figure represent hybridization between pBR322 plasmid and λ phage DNA sequences in the probes and on the filters. It is not surprising that each human transforming gene that hybridized with one also hybridized with both v-H-ras and v-K-ras because these v-onc genes share sequence homology and encode immunologically and structurally related proteins (22).

To explore further the homology between these genes, we hybridized each v-ras gene separately under conditions of high stringency to Southern blotted DNAs of the T24, SK-N-SH. and Calu-1 transforming genes and to pBR322 clones containing v-H-ras and v-K-ras (Fig. 2). As expected, v-H-ras hybridized well to the T24 transforming gene (Fig. 2A, lane c) and to the normal allele of this gene (Fig. 2A, lane d). The v-H-ras probe hybridized only weakly to a 3.0-kbp EcoRI restriction endonuclease fragment of the Calu-1 transforming gene (Fig. 2A, lane e) and to two EcoRI DNA fragments of the SK-N-SH transforming gene (Fig. 2A, lane g). In contrast, the v-K-ras probe was most closely related to the Calu-1 transforming gene, hybridizing to 3.1-, 3.0-, and 2.4-kbp EcoRI DNA fragments of this gene (Fig. 2B, lanes I and m). Longer autoradiography of the filter showed weak hybridization between v-K-ras and the T24 transforming gene (Fig. 2C, lanes j and k) and the two EcoRI fragments that comprise the SK-N-SH transforming gene (Fig. 2C, lane n).

In summary, all three human transforming genes shared homology to the v-ras genes. The T24 transforming gene was closest to v-H-ras, the Calu-1 transforming gene was closest to v-K-ras, and the SK-N-SH transforming gene was more distantly related to the v-ras genes.

The Lung and Colon Carcinoma Transforming Gene Is a Human Homolog of v-K-ras. DNAs from normal and transformed NIH 3T3 cells and from human cells were cleaved with restriction endonuclease *EcoRI* and subjected to gel electrophoresis and filter-blot hybridization, with the pKBE-2 clone

^{*}SV, simian virus; ARVT, avian reticuloendotheliosis virus T: SKV, Sloan-Kettering virus; Mu, murine.

Avian.

[‡]Personal communication.

Mammalian.

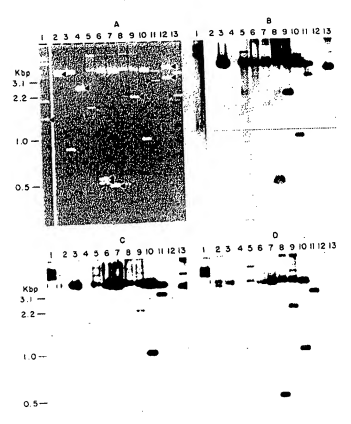


FIG. 1. Southern filter hybridization of three human transforming genes to v-onc DNA sequences. Molecular clones of retroviral oncogenes were digested with restriction endonucleases to separate oncogenic sequences from plasmid or bacteriophage DNA vectors. Digests were subjected to electrophoresis through 1% agarose gels, and the DNAs were transferred from gels to nitrocellulose filter papers (14). Filters were hybridized with cloned human transforming gene DNAs, which had been labeled with 32P by nick translation, and filters were subsequently washed under nonstringent conditions. (A) Ethidium bromide stain of a gel prior to filter transfer (arrows denote restriction endonuclease fragments containing v-onc sequences). (B-D) Filter hybridizations with ³²P-labeled pT24 DNA (B), ³²P-labeled pLC3 (C), and 32P-labeled ANPS-1-1-1(D). Lanes show the v-onc DNA restriction digests: 1, v-fps Kpn I digest of pRCII-1B; 2, v-yes Sst I purified insert from AY73-11A; 3, v-rel EcoRI digest of prel; 4, v-ski Xho I purified insert from pvski-1; 5, v-abl HindIII/Sst I digest of pABsub3; 6, v-fes Pst I digest of pGA-FeSV; 7, v-mos Pst I digest of pmos-1; 8, v-H-ras EcoRI digest of pBS-9: 9, v-H-ras EcoRI/BamHI digest of pHB-11; 10, v-K-ras EcoRI digest of pHiHi-3; 11, v-K-ras BamHI/EcoRI digest of pKBE-2; 12, v-fms Kpn I purified inserts from λ SM-FeSV; 13, v-sis EcoRI/Sal I digest of pvsis.

of v-K-ras (22) used as the 32 P-labeled hybridization probe (Fig. 3). NIH 3T3 cells transformed with DNA from the lung and colon carcinoma cells (Fig. 3, lanes 1, 2, and 3) contain K-ras-related sequences not endogenous to NIH 3T3 (Fig. 3, lane 4). The newly acquired K-ras-related EcoRI fragments in these transformed cells comigrated with v-K-ras-related EcoRI fragments prominent in human DNA (Fig. 3, lane 5). These EcoRI fragments are 2.4, 3.0, 3.1. and \approx 6.7 kbp in size. Only one high molecular weight K-ras-related EcoRI fragment in human DNA was not transferred to NIH 3T3 cells. Similar results were observed by Der et al. (5) in NIH 3T3 cells transformed with DNA from Lx-1, indicating that the same human K-ras homolog is the transforming gene of these cells.

v-K-ras Homologous Regions of the Calu-1 Transforming Gene. A large portion (26 kbp) of the transforming gene of the

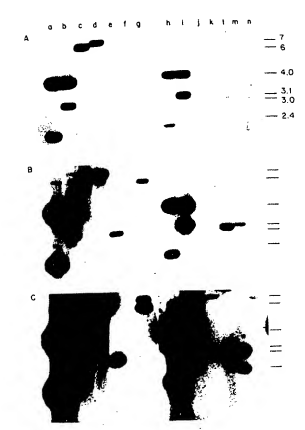


FIG. 2. Hybridization of v-ras sequence probes to filter-blotted human transforming gene DNAs. Cloned human transforming gene and v-ras gene DNAs were cleaved with restriction endonucleases, and duplicate aliquots were subjected to 1% agarose gel electrophoresis and Southern filter blotting. The filters were hybridized with either 12 P-labeled pHB-11 v-H-ras (lanes a-g) or 32 P-labeled pKBE-2 v-K-ras (lanes h-n). The filters were washed under stringent conditions. Autoradiographic exposures were for 2 hr (A), 12 hr (B), and 72 hr (C). Lanes: a and h, EcoRI/BamHI pHB-11 $(0.1 \mu g)$; b and i, EcoRI/BamHI pKBE-2 $2(0.1 \mu g)$; c and j, $2(0.1 \mu g)$; d and k, $2(0.1 \mu g)$; c and j, $2(0.1 \mu g)$; f and m, $2(0.1 \mu g)$; g and n, $2(0.1 \mu g)$;

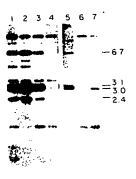
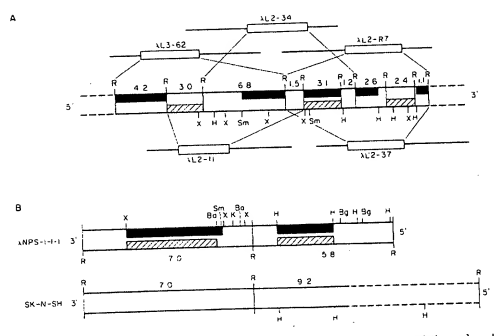


Fig. 3. Identification of lung and colon carcinoma transforming genes as K-ras homologs. Six micrograms of EcoRI-digested cellular DNA was electrophoresed through 1% agarose gels and subsequently blotted to nitrocellulose. The filters were hybridized and washed under stringent conditions with $^{32}\text{P-labeled}$ pKBE-2 (v-K-ras) as probe. Lanes 1, NIH 3T3 transformed with SK-CO-1 DNA; 2, NIH 3T3 transformed with SK-LU-1 DNA; 3, NIH 3T3 transformed with Calu-1 DNA; 4, NIH 3T3; 5, T24; 6, NIH 3T3 with 50 pg of EcoRI-cleaved $\lambda\text{L2-R7}$ DNA; 7, NIH 3T3 with 50 pg of EcoRI-cleaved $\lambda\text{L2-11}$ DNA. For structure of λ clones, see Fig. 4A. Size markers are in kbp.



human lung carcinoma cell line Calu-1 was cloned into partially overlapping A Charon 4A phages as described. A composite restriction endonuclease map for this gene is shown in Fig. 4A. Three separate regions of homology to v-K-ras were determined by hybridization analysis, comprising 3.0-, 3.1-, and 2.4kbp EcoRI fragments (see Fig. 2, lanes I and m). All three v-K-ras-related EcoRI fragments and a fourth 6.7-kbp EcoRI fragment, which has not been cloned yet, were present in all NIH 3T3 cells transformed with DNA from various lung and colon carcinoma cell lines (Fig. 3, lanes 1-3, 6, and 7). These Kirsten homologous regions do not arise by tandem gene duplications because they hybridized to discrete regions of the cloned v-K-ras gene (data not shown). Indeed, hybridization with specific v-K-ras DNA fragments allowed us to make an unambiguous assignment of the direction of transcription (see Fig. 4A). In contrast to the small T24 transforming gene, which is entirely contained on a 2.9-kbp Sac I fragment (25), the transforming gene of Calu-1 is probably greater than 30 kbp.

The SK-N-SH Neuroblastoma Transforming Gene Is a New Member of the ras Gene Family. Although the SK-N-SH neuroblastoma transforming gene was weakly homologous to both v-H-ras and v-K-ras, we reasoned that it may encode a protein structurally and serologically related to Harvey and Kirsten ras gene product. We tested this possibility by using a broadly reactive monoclonal antibody against ras-encoded protein to immunoprecipitate [35S]methionine-labeled extracts from three independently derived NIH 3T3 transformants containing the SK-N-SH transforming gene. Immune precipitates from these cells and from NIH 3T3 transformed by Harvey sarcoma virus unintegrated DNA. NIH 3T3 transformed by DNAs from human lung and colon carcinoma cells, and NIH 3T3 itself were analyzed by NaDodSO₄/polyacrylamide gel electrophoresis (Fig.

5). A protein with an apparent $M_{\rm r}$ of 19,000 was seen in immunoprecipitates of v-H-ras-transformed NIH 3T3 (Fig. 5, lane 2) but not in NIH 3T3 controls (Fig. 5, lane 1). A similarly migrating protein was seen in NIH 3T3 cells transformed with either

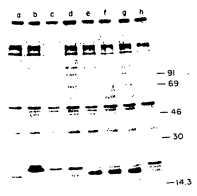


Fig. 5. ras-Related proteins in NIH 3T3 transformed cells. Cells (5 \times 10⁵) were labeled for 18 hr with 80 μ Ci of (35S)methionine and extracted with nonionic detergents, and cleared lysates were used for immunoprecipitation with rat anti-ras p21 monoclonal antibody Y13-259 (16). Immunocomplexes were collected onto S. aureus protein A, dissolved and boiled in NaDodSO₄ sample buffer, and analyzed by ADOdSO₄/polyacrylamide gel electrophoresis on a 12.5% polyacrylamide gel. Proteins were visualized by fluorography. Lanes: a, NIH 3T3; b, HT14B; c and d, two NIH 3T3 lines independently transformed with Sk-N-SH DNA: e-g, three NIH 3T3 lines independently transformed with SK-N-SH DNA: h, one NIH 3T3 line transformed with SK-CO-1 DNA. A ¹⁴C-labeled protein mixture (Amersham) provided M_r standards (shown \times 10⁻³).

Calu-1 or SK-CO-1 DNA (Fig. 5, lanes 3, 4, and 8). A uniquely migrating protein with an apparent Mr of 17,500 was seen in each NIH 3T3 transformant containing the SK-N-SH transforming gene (Fig. 5, lanes 5-7). This protein had an isoelectric point similar to that found for the v-H-ras-encoded protein (data not shown). Our findings indicate that the SK-N-SH neuroblastoma transforming gene is another member of the ras gene family.

We exploited the homology between the SK-N-SH transforming gene and the v-ras genes to determine its direction of transcription and approximate regions of homology (see Fig. 4B).

DISCUSSION

Three different human transforming genes that can be detected by the NIH 3T3 transformation assay are members of the ras gene family. The transforming gene of a bladder carcinoma cell line (T24) is a human homolog of the v-H-ras gene (5, 10, 11). The transforming gene of Lx-1, a human lung carcinoma cell line, is a human homolog of v-K-ras (5). Comparison of the work of Der et al. (5) with ours indicates the presence of the same transforming gene in two lung carcinoma cell lines (SK-LU-1 and Calu-1) and in one colon carcinoma cell line (SK-CO-1). This same gene is also detectable by DNA transfer in human lung and colon tumors maintained in nude mice (unpublished data) and in the colon carcinoma cell line SW480 (3, 9). The transforming gene of a human neuroblastoma cell line (SK-N-SH) is related to (but distinct from) the homologs of the v-Hras and v-K-ras genes and represents a third branch within the ras gene family. Each branch may have more recent evolutionary offshoots. Thus, Chang et al. (34) reported two human homologs of v-H-ras (H-ras-1 and -2) and two homologs of v-K-ras (K-ras-1 and -2). A comparison of restriction endonuclease maps for these genes with the three human transforming genes we have isolated indicates that the T24 bladder carcinoma-transforming gene is H-ras-1, the lung and colon carcinoma-transforming gene is probably K-ras-2, and the SK-N-SH neuroblastoma-transforming gene is a heretofore uncharacterized gene. We propose calling the human transforming gene of SK-N-SH the N-ras-1 gene.

It is of considerable interest that a wide variety of tumor cells contain activated ras genes. detectable by gene transfer into NIH 3T3 cells. Several factors possibly contribute: ras transforming genes may be more readily detected than other transforming genes by the NIH 3T3 focus assay; ras genes may be easily activated by mutation; and ras genes may have critical cellular functions in a wide variety of cell types. The function of the ras gene products is not known nor is it known whether they perform physiologically distinguishable roles. However, it is known that an altered amino acid sequence is responsible for the activation of the H-ras-1 gene of T24 (25, 35, 36), and we speculate that alteration in the ras gene products may be a common step in many forms of human cancer.

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